

Alfa Aesar[®]

A Johnson Matthey Company

**Research
Chemicals,
Metals and
Materials**

New Products Addendum

March 2010

Also Available from Alfa Aesar

RESEARCH CHEMICALS, METALS AND MATERIALS CATALOG

The Research Chemicals, Metals and Materials catalog offers more than 30,000 products. The complete range of organics, inorganics, metals and materials includes over 3,000 new products.

FINE CHEMICALS & CATALYSTS BROCHURE

The 14-page brochure outlines Alfa Aesar's key capabilities as a supplier and manufacturer of fine chemicals, including organics, high purity inorganics and precious metal catalysts.

BORONIC ACIDS BROCHURE

This 56-page technical document outlines the chemical properties and highlights the main synthetic uses of boronic acids and related chemicals.

HIGH PURITY INORGANICS CATALOG

This catalog lists over 1,300 high purity inorganic materials, including base metal compounds, rare earth compounds and ultra dry materials for air and moisture sensitive applications.

ANALYTICAL PRODUCTS CATALOG

The 256-page catalog highlights Alfa Aesar's Specpure® range of analytical standards and lists hundreds of other products including HPLC solvents, high purity acids and bases, NMR and specialty solvents, MS standards, labware and more.

HIGH PURITY MATERIALS FOR PHOTOVOLTAICS BROCHURE

The 8-page brochure reviews Alfa Aesar's range of High Purity Photovoltaic (PV) compounds and elements specifically for the Solar Industry, including silicon, CIGS and CdTe.

PRECIOUS METAL COMPOUNDS AND CATALYSTS CATALOG

This 80-page publication lists a full range of precious metal chemicals and catalysts, including FibreCat™ anchored homogeneous catalysts and HiSPEC™ fuel cell catalysts.

PLATINUM LABWARE CATALOG

The Platinum Labware Catalog includes a large selection of crucibles, molds, dishes, micro chemical apparatus and utensils.

SPECTROFLUX® ANALYTICAL FLUXES BROCHURE

Our 10-page color Spectroflux Information Package illustrates the wide range of alkali metal borate fusion fluxes available for use in AA/ICP analytical techniques.

NANOPARTICLES AND DISPERSIONS BROCHURE

This 8-page brochure describes the manufacture and full range of both metal oxide powders and dispersions manufactured by Nanophase and distributed solely by Alfa Aesar.

HIGH PURITY METALS & MATERIALS CATALOG

The High Purity Metals and Materials Catalog highlights Alfa Aesar's entire range of metals and alloys from aluminum to zirconium, featuring purities up to 99.9999%. Products are offered in a broad variety of forms, including wires, foils, shots, targets, powders, thermocouple wires, and many more.

FUEL CELL CATALYSTS & COMPONENTS BROCHURE

This brochure describes Alfa Aesar's offering of high quality Johnson Matthey fuel cell products, including HiSPEC™ brand precious metal fuel cell catalysts, HiFUEL™ brand base metal fuel cell catalysts, plus a range of electrodes and assemblies.

SPECIALTY SILVER COMPOUNDS BROCHURE

The 8-page brochure describes Alfa Aesar's expertise in producing silver compounds up to 99.999% (metals basis) purity and in batches as large as 200kg. Seventy-five standard products are listed and Alfa Aesar can also manufacture to meet custom specifications.

Go to www.alfa.com for a full selection of literature available in downloadable PDF format.



How to Order

TELEPHONE

Catalog Sales

(800) 343-0660 or (978) 521-6300

Catalog Sales Hours:

Monday–Friday, 8:00 AM–7:00 PM EST

Specialty and Bulk Sales

(888) 343-8025 or (978) 521-6401

Monday–Friday, 8:00 AM–5:00 PM EST

Technical Service

(800) 343-7276 or (978) 521-6405

Monday–Friday, 8:00 AM–5:00 PM EST

Please have the following information available when placing an order:

- Your customer number (if applicable).
- Billing and shipping addresses.
- Product stock number, size, and quantity for each item you wish to order.
- Your purchase order number or government/corporate credit card number and expiration date.

FAX

Catalog Sales

(800) 322-4757 or (978) 521-6350

Specialty and Bulk Sales

(978) 521-6366

MAIL

Alfa Aesar

26 Parkridge Road
Ward Hill, MA 01835-6904

INTERNET

www.alfa.com

Our web catalog features up-to-date prices and a user-friendly e-commerce system.

EMAIL

info@alfa.com

for catalog product orders and general inquiries

tech@alfa.com

for technical inquiries

specialquotes@alfa.com

for bulk and specialty inquiries

We do not require a confirming purchase order. If you choose to send one, please mark it clearly, “CONFIRMING ORDER” to avoid possible duplication. There is no minimum order. We welcome all orders, regardless of size.

Alfa Aesar is ISO 9001:2000 Certified

General Information

ORDERING

Customer Service Representatives are available weekdays from 8:00AM to 7:00PM EST. When you call (800)343-0660 to place an order, we will work with you to learn your specific needs. There is no minimum order. All orders are accepted, regardless of size.

PRICING

All prices are listed in U.S. dollars. Every effort is made to hold prices firm during the life of the catalog. However, Prices Are Subject To Change Without Notice. Most current pricing may be found at our website: www.alfa.com. In cases where the selling price has changed significantly, we will contact you prior to filling your mailed or faxed order. Our payment terms are net 30 days of invoice.

SHIPPING

Whenever possible, we will ship products by the method specified on your order. Ground delivery is standard within the contiguous United States. All orders are shipped FOB seller's shipping point.

TECHNICAL SERVICE

At your request, we will furnish technical assistance and information with respect to our products. Our Technical Service Representatives are trained in specific product lines to answer your questions regarding applications, specifications, product properties and handling. Call (800) 343-7276.

MATERIAL SAFETY DATA SHEETS

Each product ordered is automatically accompanied by a Material Safety Data Sheet (MSDS) in compliance with OSHA. If one is not immediately available, a copy will be sent via mail as soon as possible. If an MSDS is needed prior to shipment of a product please call (800) 343-0660 or visit our website at www.alfa.com.

CERTIFICATES OF ANALYSIS

Certificates of analysis are available online at www.alfa.com or by calling (800)343-0660. All Premion®, Puratronic®, REacton®, and Specpure® brand products are automatically accompanied by a batch-specific Certificate of Analysis.

RETURN SHIPMENTS

Some materials are not returnable. Returned shipments cannot be accepted unless prior arrangements have been made. Requests for return authorization must be made within 30 days of your receiving the materials. A 20% restocking fee is charged on authorized returns of standard catalog items, subject to a minimum charge of \$20.00 per order.

TERMS OF SALE

Full details of Terms and Conditions are listed on our website (www.alfa.com). For health and safety reasons, we shall not supply chemicals to private individuals or deliver to residential addresses. Orders will be accepted from bona fide business customers only.

NEW CUSTOMERS

We welcome new customers and setting up an account with Alfa Aesar is easy. Just call (800) 343-0660 and a customer service representative will assist you.

Key to Chemical Listings

1	2	3	10	11	4	5
B24303	Benzhydrylamine, 97% Δ				5g 28.40	
6	[Aminodiphenylmethane]				25g 58.80	
	[91-00-9], (C ₁₃ H ₁₃) ₂ CHNH ₂ , F.W. 183.25, m.p. 11-13°, b.p. 295°, f.p. >110° (230°F), d. 1.064, n _D ²⁰ 1.5950, Merck 14,1076, Solubility: Slightly soluble in water, UN2735, MDL MFCD00008059, †	13	8	16	100g 195.00	
L02388	Benzhydrylamine hydrochloride, 97%					
7	[Aminodiphenylmethane hydrochloride]					
	[5267-34-5], C ₁₃ H ₁₃ N·HCl, F.W. 249.72, m.p. 293-295°, Merck 14,1076, EINECS 226-084-0, BRN 3914818, MDL MFCD00013023				10g 19.50	
12					50g 79.50	
14	Can be used as a protected form of ammonia in a practical synthesis of azetidine, suitable for use on a multi-kilo scale: Synth. Commun., 18, 205 (1988):			15		
44459	Benzhydrylamine hydrochloride, resin bound				5g 45.50	
9	[BHA·HCl resin] Powder, MDL MFCD03456188, Note: Poly(styrene-divinylbenzene), 1% cross-linked, 100-200 mesh base resin, loading range 0.5-1.5mmol/g				25g 183.00	

- | | |
|---|---|
| 1 Stock number | 10 UN Dangerous Goods number |
| 2 Product name and grade | 11 Denotes substance is listed in Toxic Substance Control Act (TSCA) inventory |
| 3 Sensitivity symbol (see abbreviations index) | 12 Hazard and storage indicators |
| 4 Unit size | 13 European Union Risk and Safety Phrase Codes |
| 5 Prices | 14 Product review with literature references |
| 6 Alternative name(s) | 15 Chemical structure |
| 7 Chemical Abstract Service (CAS) number | 16 EINECS (European Inventory of Existing Chemical Substances) Number |
| 8 Molecular formula | |
| 9 Physical form | |

F.W.	Formula weight
m.p.	Melting point
b.p.	Boiling point
f.p.	Flash point
d.	Density in g/ml
n_D²⁰	Refractive index
[α]_D²⁰	Optical rotation

Merck	Merck Index Number
BRN	Beilstein Registry number
Fieser	Fieser's Reagents reference
RTECS	Registry of Toxic Effects of Chemical Substances reference
MDL	MDL number

Bulk & Specialty Capabilities

The expert technical and commercial personnel in our Bulk and Specialty Sales group can aid in your search for the ideal bulk, special or custom manufactured product.

MANUFACTURING

Alfa Aesar's combined resources bring you a broader range of unique building blocks and other novel compounds. Over 100 production chemists employ a variety of techniques in the development of new products, many of which are otherwise unavailable commercially.

Alfa Aesar's production operations offer a single source for fine chemicals, from research to pilot scale. Full-scale production for many products is available through various Johnson Matthey plants worldwide.

CUSTOM SYNTHESIS

We offer flexible custom manufacturing services with the assurance of quality and confidentiality. We can deliver the chemical you need in sizes for research, pilot-scale and full-scale production applications.

CUSTOMER SERVICE

Our dedicated scientific and commercial teams offer full service from production to delivery. Most products are stocked in catalog pack sizes and the majority are available from stock in semi-bulk and bulk quantities as well. All specialty and bulk products are shipped with a batch specific certificate of analysis and material safety data sheet. Because we understand that specific packaging is often important, we offer custom packaging and labeling to meet your requirements.



Abbreviations and Codes

The following abbreviations are used throughout our listing of products.

AAS	Atomic absorption spectrometry	N	Normality of solution
AES	Atomic emission spectrometry	n_D^{20}	Refractive index for the sodium D line at 20 °C (or temperature indicated)
ACS	Chemicals meeting the specifications outlined by the American Chemical Society	nm	Nanometer
APS	Average particle size		New product
anhy	Anhydrous	NMR	Nuclear magnetic resonance
Å	Angstrom	OD	Outer diameter
≈	Approximately	optical gr.	Suitable for optical applications
approx.	Approximately	pc(s)	Piece(s)
Atm	Atmospheres	pH	Value taken to represent the acidity or alkalinity of an aqueous solution
b.p.	Boiling point in °C at 760mm pressure, unless otherwise specified	POR	Price on request
(c)	Contained weight of active material	ppb	Parts per billion
°C	Celsius	ppm	Parts per million
cc	Cubic centimeter	prec.	Precipitated
cm	Centimeter	Primary	Analytical reagent of exceptional purity, for standardizing volumetric solutions and preparing reference standards
cont.	Contained	Standard	
cP	Centipoise	PTFE	Poly(tetrafluoroethylene)
cS	Centistoke	Purified	A grade of higher quality than technical, often used where there are no official standards
d.	Density	P.T.	Passes test
dec.	Decomposes	P.V.	Pore volume
dia.	Diameter	Reagent	Reagent grade
ea.	Each	REM	Rare earth metal
eV	Electron volt	(REO)	Rare earth oxide base - content of specific rare earth element in comparison to total rare earths present
°F	Fahrenheit	S.A.	Surface area
f.p.	Flash point	Sp.Gr.	Specific gravity
FSSS	Fisher sub-sieve sizer	Sp.Rot.	Specific rotation
F.W.	Formula weight	subl.	Sublimes
g	Gram	Tc	Critical temperature
g/l	Grams per liter (gas density)	TLC	Thin-layer chromatography
GLC	Suitable for use in gas liquid chromatography	TSCA	Toxic Substance Control Act
GC	Gas chromatography	UN	Hazardous material transportation identification number
HPLC	High-performance liquid chromatography	λ	Wavelength in nanometers
ICP	Inductively Coupled Plasma	wt	Weight
ID	Inner diameter	w/w	Weight/weight
in	Inch	w/v	Weight/volume
incl	Includes	XRD	X-ray diffraction
IR	Infrared	△	Air sensitive
J/mol·K	Joule(s) per mole Kelvin	▮	Moisture sensitive
kg	Kilogram	■	Hygroscopic
L or l	Liter	▲	Light sensitive
lb	Pound	>	Greater than
m	Meter	≥	Greater than or equal to
M	Molarity of solution	<	Less than
mg	Milligram	≤	Less than or equal to
μ	Micro	[]	Numbers in brackets after the chemical description indicate the Chemical Abstract Service Registry Number
μg	Microgram	- mesh #	90% particles pass through screen having a given mesh size
meq	Milliequivalent	+ mesh #	90% particles are retained by a screen having a given mesh size
Merck	The Merck Index	†	Denotes substance is listed in Toxic Substance Control Act (TSCA) inventory
μm	Micrometer (micron)		
max	Maximum		
micron	Micrometer		
mm	Millimeter		
m.p.	Melting point		
mol	Mole		
mmol	Millimole		
M.W.	Molecular weight		
Mw	Weighted averaged molecular weight		
Mn	Number averaged molecular weight		
Mw/Mn	Monodispersity value		

Section 1 Compounds

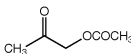
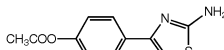
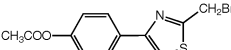
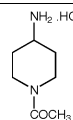
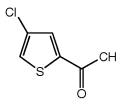
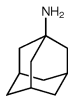
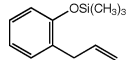
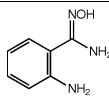
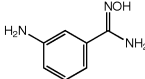
Alphabetical listing of compounds. 4

This section is an alphabetical listing of new compounds added since the publication of the 2008-09 Alfa Aesar Catalog. We have, where practical, used the systematic nomenclature adopted by the International Union of Pure and Applied Chemistry (IUPAC). Some products, however, are listed under their Chemical Abstract, trivial or common name. Cross-references have been provided to assist you in finding the product. Many product listings include extended application notes and related literature references.

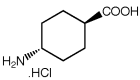
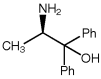
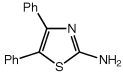
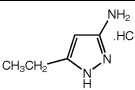
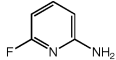
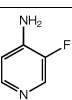
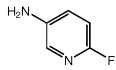
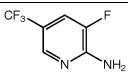
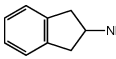
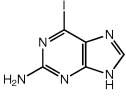
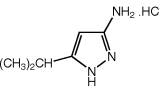
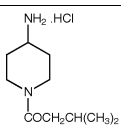
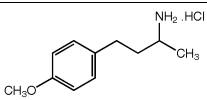
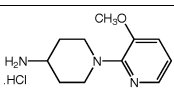
For most products, the Chemical Abstracts Service Registry Number (CAS) is indicated. To assist users of our catalog, representative physical constants and solubilities (mainly literature values) are included with the listings. These figures are not specifications and are supplied for informational purposes only.

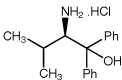
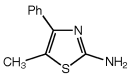
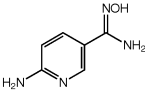
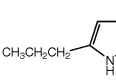
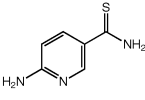
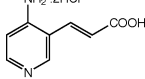
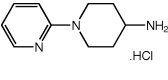
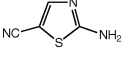
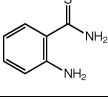
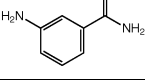
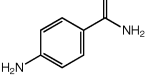
Air, moisture and/or hygroscopic sensitivities, where known, are indicated to alert our customers that special precautions are needed in handling a particular compound. Note that no attempt is made to indicate the degree of air, moisture, or hygroscopic sensitivity in any given case. For example, the term "moisture sensitive" is applied to all degrees of reactivity from hygroscopic to spontaneously flammable. The terms "air sensitive", "light sensitive" and "hygroscopic" are used in a similarly broad sense.

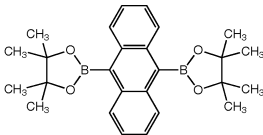
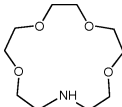
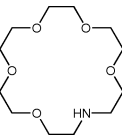
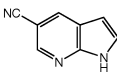
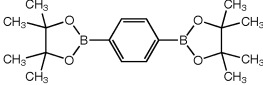
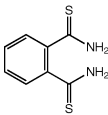
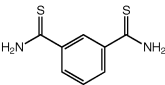
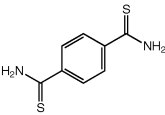
For a complete list of our chemicals, metals and materials, please visit our online catalog at www.alfa.com. The site lists all products with up-to-date price and availability information.

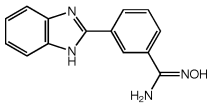
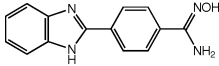
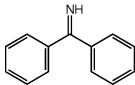
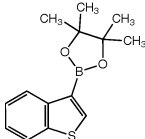
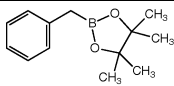
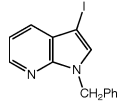
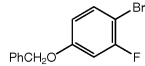
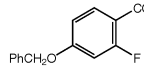
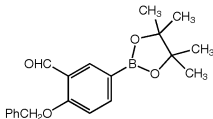
Stock #	Description		Size	\$
H31346	Acetoxyacetone, 97% [Acetonyl acetate, 1-Acetoxy-2-propanone] [592-20-1], C ₅ H ₈ O ₃ , F.W. 116.12, b.p. 174-176°, f.p. 71°(160°F), d. 1.075, EINECS 209-746-3, †		10ml 50ml	64.60 241.00
H51858	4-(4-Acetoxyphenyl)-2-aminothiazole, 97% C ₁₁ H ₁₀ N ₂ O ₂ S, F.W. 234.27 ☒ R:22-36/37/38, S:26-36/37		1g 5g	67.00 249.00
H51822	4-(4-Acetoxyphenyl)-2-(bromomethyl)thiazole, 97% C ₁₂ H ₁₀ BrNO ₂ S, F.W. 312.18, UN3261 ☒ R:34, S:26-36/37/39-45		1g 5g	67.00 249.00
H30389	1-Acetyl-4-aminopiperidine hydrochloride, 97% △ [1-Acetyl-4-piperidinamine hydrochloride] [214147-48-5], C ₇ H ₁₄ N ₂ O·HCl, F.W. 178.66, m.p. 231-234° dec. ☒ R:36/37/38, S:26-37		250mg 1g	70.10 253.00
H30384	2-Acetyl-4-chlorothiophene, 98+% [34730-20-6], C ₄ H ₃ ClOS, F.W. 160.62, d. 1.334, UN2810, MDL MFCD00082791 ☒ R:20/21/22, S:36/37		1g 5g 25g	29.30 101.00 383.00
H30076	1-Adamantanamine, 98% [768-94-5], C ₁₀ H ₁₇ N, F.W. 151.25, m.p. ca 200° subl., EINECS 212-201-2, RTECS YD1925000, † ☒ R:22-36/37/38, S:26-36/37		5g 25g 100g	42.60 136.00 412.00
H32601	Agarose beads 4% B [9012-36-6], b.p. 100°, f.p. 38°(100°F), d. 1.040, UN1170, † R:10		1L	595.00
H32305	Agarose beads 4% B-CL (Cross-linked) [61970-08-9], b.p. 100°, f.p. 38°(100°F), d. 1.040, UN1170 R:10		1L	665.00
H32491	Agarose beads 6% B [9012-36-6], b.p. 100°, f.p. 38°(100°F), d. 1.040, UN1170, † R:10		1L	890.00
H32568	Agarose beads 6% B-CL (Cross-linked) [62610-50-8], b.p. 100°, f.p. 38°(100°F), d. 1.040, UN1170 R:10		1L	745.00
H51170	Allylmagnesium chloride, 1M in MeTHF △ ▯ [2622-05-1], CH ₂ =CHCH ₂ MgCl, F.W. 100.83, UN3399, † ☒ R:11-14/15-19-34, S:8-16-26-36/37/39-43-45		100ml 500ml	42.00 158.00
H30905	(2-Allylphenoxy)trimethylsilane, 98% [18042-43-8], C ₁₂ H ₁₈ OSi, F.W. 206.36, b.p. 85°/5mm, d. 0.925 ☒ R:36/38, S:26-37		5g 25g 100g	59.80 239.00 755.00
45586	Aluminum oxide, NanoArc® AL-2220, 30% in mineral spirits, colloidal dispersion with dispersant [1344-28-1], Al ₂ O ₃ , F.W. 101.96, Application(s): Aluminum oxide dispersed in mineral spirits suitable for incorporation in non-polar organic based coatings and formulations where high transparency is important, UN1993, EINECS 215-691-6, MDL MFCD00003424, Note: Sold in collaboration with Nanophase Technologies Corporation R:10		100g 500g	27.50 100.00
45482	Aluminum oxide, alpha-phase, 99.98% (metals basis) [1344-28-1], Al ₂ O ₃ , F.W. 101.96, m.p. 2045°, b.p. 2980°, n _D ²⁰ 1.765, Merck 14,356, EINECS 215-691-6, MDL MFCD00003424, †		25g 100g 500g	15.50 41.00 98.70
H32475	Amberlite® IRA-96, ion-exchange resin [39409-19-3]		250g 1kg	40.60 115.00
45591	Amberlite® IRN-78, ion exchange resin, nuclear grade [11128-95-3], Styrene-DVB; Strongly basic anion exchange resin, Merck 14,382, †		500g 2kg	23.80 47.30
H51782	2-Aminobenzamidoxime, 97% [16348-49-5], C ₇ H ₈ N ₂ O, F.W. 151.17, m.p. 71-75°, MDL MFCD00492723 ☒ R:36/37/38, S:26-37		1g 5g	57.60 214.00
H51773	3-Aminobenzamidoxime, 97% [100524-07-0], C ₇ H ₈ N ₂ O, F.W. 151.17, m.p. 101-105°, MDL MFCD08061150 ☒ R:36/37/38, S:26-37		1g 5g	67.20 250.00

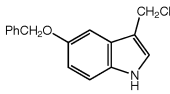
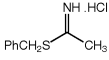
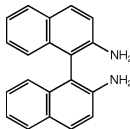
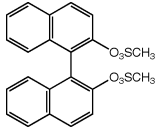
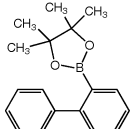
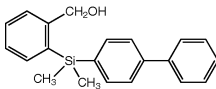
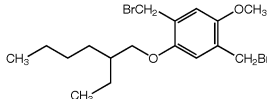
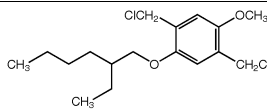
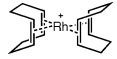
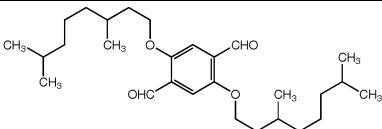
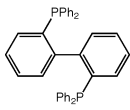
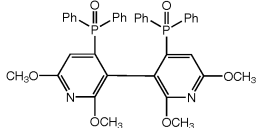
Stock #	Description		Size	\$
H51788	4-Aminobenzamidoxime, 97% [277319-62-7], C ₇ H ₉ N ₃ O, F.W. 151.17, m.p. 168-170°, MDL MFCD08061151 ✗ R:36/37/38, S:26-37		1g 5g	63.40 236.00
H51666	4-Aminobenzeneboronic acid pinacol ester, 98% [214360-73-3], C ₁₂ H ₁₅ BN ₂ O ₅ , F.W. 219.09, MDL MFCD02093721		1g 5g	59.90 215.00
H51027	3-Aminobiphenyl hydrochloride [3-Biphenylamine hydrochloride, 3-Phenylaniline hydrochloride] [2113-55-5], C ₁₂ H ₁₁ N·HCl, F.W. 205.69, MDL MFCD00068358 ✗ R:22-40, S:36/37		1g 5g	107.00 481.00
H30538	(S)-(+)-3-Amino-1-Boc-piperidine, 97% △ [(S)-Boc-3-piperidinamine, tert-Butyl (R)-3-aminopiperidine-1-carboxylate] [625471-18-3], C ₁₀ H ₂₀ N ₂ O ₂ , F.W. 200.28, UN2735 ✗ R:34, S:26-36/37/39-45		250mg 1g 5g	57.00 158.00 528.00
H30779	4-Amino-1-Boc-piperidine, 97% △ [1-Boc-4-piperidinamine, tert-Butyl 4-aminopiperidine-1-carboxylate] [87120-72-7], C ₁₀ H ₂₀ N ₂ O ₂ , F.W. 200.28, m.p. 44-48°, UN3259 ✗ R:34, S:26-36/37/39-45		1g 5g	58.50 148.00
H31614	3-Amino-5-bromo-2-chloropyridine, 97% [588729-99-1], C ₅ H ₄ BrClN ₂ , F.W. 207.46, m.p. 132-135°, MDL MFCD02682092 ✗ R:20/21/22-36/37/38, S:26-36/37		1g 5g	53.70 180.00
H31500	2-Amino-4-bromopyridine, 97% [84249-14-9], C ₅ H ₄ BrN ₂ , F.W. 173.01, m.p. 144-147° ✗ R:20/21/22-36/37/38, S:26-36/37		250mg 1g	75.10 210.00
H32526	3-Amino-4-bromopyridine, 96% [239137-39-4], C ₅ H ₄ BrN ₂ , F.W. 173.01, b.p. 46-49°/3mm, MDL MFCD04112575 ✗ R:20/21/22-36/38, S:26-36/37		250mg 1g	58.50 162.00
H31514	2-Amino-5-bromothiazole hydrobromide, 97% [61296-22-8], C ₃ H ₃ BrN ₂ S·HBr, F.W. 303.95, m.p. ca 165° dec., EINECS 262-699-0, RTECS XJ1262200, MDL MFCD00012712 ✗ R:36/37/38, S:26-37		1g 5g 25g	22.60 64.60 195.00
H32991	2-Amino-3-bromo-5-(trifluoromethyl)pyridine, 97% [79456-30-7], C ₆ H ₄ BrF ₃ N ₂ , F.W. 241.01, m.p. 98-101°, MDL MFCD07375382 ✗ R:20/21/22-36/37/38, S:26-36/37		250mg 1g	27.70 76.50
H30531	2-Amino-5-chlorobenzyl alcohol, 98% [37585-25-4], C ₇ H ₈ ClNO, F.W. 157.60, m.p. 107-109° ✗ R:36/37/38, S:26-37		5g 25g	76.00 252.00
H30135	2-Amino-5-chloro-3-fluoropyridine, 98% [5-Chloro-3-fluoro-2-pyridinamine] [246847-98-3], C ₅ H ₄ ClFN ₂ , F.W. 146.55, m.p. 95-97° ✗ R:20/21/22-36/37/38, S:26-36/37		250mg 1g 5g	67.80 189.00 628.00
H51085	5-Amino-1-(4-chlorophenyl)-3-methyl-1H-pyrazole, 97% △ [40401-39-6], C ₁₀ H ₁₀ ClN ₃ , F.W. 207.66, EINECS 254-907-3, MDL MFCD03478180, † ✗ R:36/37/38, S:26-37		1g	144.00
H30166	2-Amino-3-chloropyridine, 95% [3-Chloro-2-pyridinamine] [39620-04-7], C ₅ H ₅ ClN ₂ , F.W. 128.56, m.p. 59-62° ✗ R:20/21/22-36/37/38, S:26-36/37		250mg 1g 5g	65.50 182.00 609.00
H31093	4-Amino-3-chloropyridine, 97% [3-Chloro-4-pyridinamine] [19798-77-7], C ₅ H ₅ ClN ₂ , F.W. 128.56, m.p. 49-53° ✗ R:20/21/22-36/37/38, S:26-36/37		1g 5g	142.00 472.00

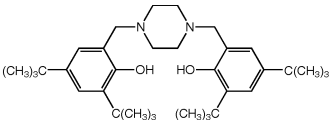
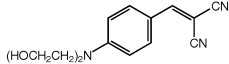
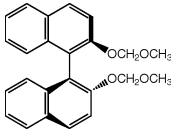
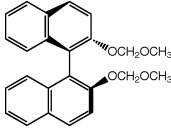
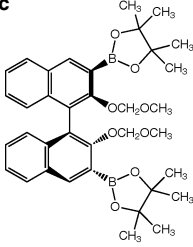
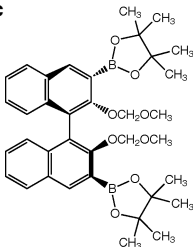
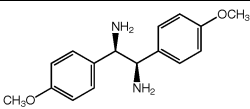
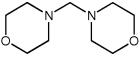
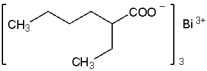
Stock #	Description		Size	\$
H31114	trans-4-Aminocyclohexanecarboxylic acid hydrochloride, 96% [27960-59-4], C ₆ H ₁₃ NO ₂ ·HCl, F.W. 179.65, m.p. >300°  R:36/37/38, S:26-37		1g 5g	63.20 210.00
H51083	(R)-(+)-2-Amino-1,1-diphenyl-1-propanol, 99% ▲ [78603-93-7], C ₁₅ H ₁₇ NO, F.W. 227.30, m.p. 102-105°, UN3259, MDL MFCD02677967  R:34, S:26-36/37/39-45		1g 5g	129.00 522.00
H51818	2-Amino-4,5-diphenylthiazole, 97% [6318-74-7], C ₁₅ H ₁₃ N ₂ S, F.W. 252.33, RTECS XJ2450000, MDL MFCD00438803  R:22-36/37/38, S:26-36/37		1g 5g	53.00 197.00
H51026	3-Amino-5-ethyl-1H-pyrazole hydrochloride C ₅ H ₇ N ₃ ·HCl, F.W. 147.61, MDL MFCD11111022  R:36/37/38, S:26-37		250mg 1g	59.10 213.00
H31970	2-Amino-3-fluoropyridine, 97% [21717-95-3], C ₅ H ₅ FN ₂ , F.W. 112.11, m.p. 39-43°, UN2811, MDL MFCD04114135  R:21-25-36/37/38, S:26-36/37-45		250mg 1g	50.40 140.00
H30629	2-Amino-6-fluoropyridine, 97% [6-Fluoro-2-pyridinamine] [1597-32-6], C ₅ H ₅ FN ₂ , F.W. 112.11, m.p. 58-62°, UN2811  R:21-25-36/37/38, S:26-36/37-45		1g 5g	85.20 284.00
H32949	4-Amino-3-fluoropyridine [2247-88-3], C ₅ H ₅ FN ₂ , F.W. 112.11, UN2811  R:21-25-36/37/38, S:26-36/37-45		1g	94.70
H30872	5-Amino-2-fluoropyridine, 97+% [6-Fluoro-3-pyridinamine] [1827-27-6], C ₅ H ₅ FN ₂ , F.W. 112.11, m.p. 88-93°, UN2811  R:21-25-36/37/38, S:26-36/37-45		1g 5g	273.00 910.00
H32532	2-Amino-3-fluoro-5-(trifluoromethyl)pyridine, 97% [852062-17-0], C ₈ H ₄ F ₄ N ₂ , F.W. 180.10, m.p. 64-68°, UN2811  R:24/25-36/37/38		250mg 1g 5g	61.40 170.00 565.00
H31023	2-Aminoindane, 97% [2975-41-9], C ₉ H ₁₁ N, F.W. 133.19, b.p. 229°, f.p. 100° (212°F), d. 1.024, UN2735, EINECS 221-021-3   R:22-34, S:26-36/37/39-45		1g 5g	35.60 122.00
H51672	2-Amino-6-iodopurine, 97% ▲ [19690-23-4], C ₅ H ₄ IN ₅ , F.W. 261.02		1g 5g	28.50 110.00
H51110	3-Amino-5-isopropyl-1H-pyrazole hydrochloride C ₈ H ₁₁ N ₃ ·HCl, F.W. 161.6, MDL MFCD11111023  R:36/37/38, S:26-37		250mg 1g	53.80 194.00
H51684	4-Amino-1-isovalerylpyperidine hydrochloride, 99% C ₁₀ H ₂₀ N ₂ O·HCl, F.W. 220.74  R:36/37/38, S:26-37		250mg 1g	60.70 218.00
H51680	3-Amino-1-(4-methoxyphenyl)butane hydrochloride, 98% C ₁₁ H ₁₇ NO·HCl, F.W. 215.72, m.p. 128-131°  R:36/37/38, S:26-37		250mg 1g	67.80 244.00
H51176	4-Amino-1-(3-methoxy-2-pyridyl)pyperidine hydrochloride C ₁₇ H ₁₇ N ₃ O·HCl, F.W. 243.73, MDL MFCD11869912  R:22-36/37/38, S:26-36/37		250mg 1g	74.30 246.00

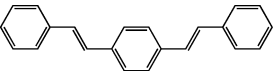
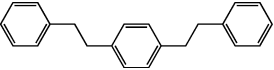
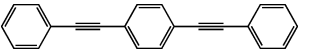
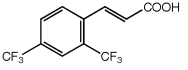
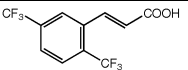
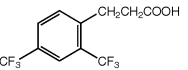
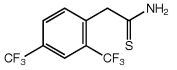
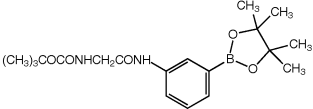
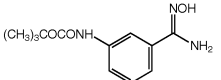
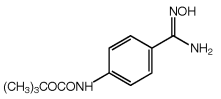
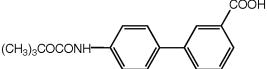
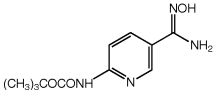
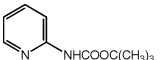
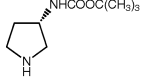
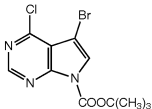
Stock #	Description		Size	\$
H51069	(R)-(+)-2-Amino-3-methyl-1,1-diphenyl-1-butanol hydrochloride, 98% C ₁₇ H ₂₁ NO·HCl, F.W. 291.85, MDL MFCD01318247  R:36/37/38, S:26-37		1g	126.00
			5g	462.00
H51825	2-Amino-5-methyl-4-phenylthiazole, 97% [30709-67-2], C ₁₀ H ₁₀ N ₂ S, F.W. 190.26, m.p. 107-112°, RTECS XJ2650000, MDL MFCD00617257  R:22-36/37/38, S:26-36/37		1g	41.00
			5g	153.00
H30935	3-Amino-1-methyl-1H-pyrazole, 97% Δ [1-Methyl-1H-pyrazol-3-amine] [1904-31-0], C ₄ H ₅ N ₃ , F.W. 97.12, b.p. 90-92°/1mm, UN2735, MDL MFCD00466340   R:22-34, S:26-36/37/39-45		1g	41.30
			5g	138.00
H51720	6-Aminonicotinamidoxime, 97% [468068-28-2], C ₆ H ₈ N ₄ O, F.W. 152.16, m.p. 192-195°  R:36/37/38, S:26-37		250mg	110.00
			1g	315.00
H51024	3-Amino-5-n-propyl-1H-pyrazole hydrochloride C ₈ H ₁₁ N ₃ ·HCl, F.W. 161.63, MDL MFCD11111020  R:36/37/38, S:26-37		250mg	54.70
			1g	197.00
H51775	2-Aminopyridine-5-thiocarboxamide, 97% [53268-33-0], C ₆ H ₆ N ₂ S, F.W. 153.20, m.p. 172-175°  R:22-36/37/38, S:26-36/37		250mg	80.60
			1g	257.00
H51665	3-(4-Amino-3-pyridyl)acrylic acid dihydrochloride C ₈ H ₈ N ₂ O ₂ ·2HCl, F.W. 237.08, MDL MFCD11870729  R:36/37/38, S:26-37		250mg	56.70
			1g	204.00
H51108	4-Amino-1-(2-pyridyl)piperidine hydrochloride [1-(2-Pyridyl)-4-piperidinamine hydrochloride] C ₁₀ H ₁₅ N ₃ ·HCl, F.W. 213.71, MDL MFCD09607833  R:36/37/38, S:26-37		250mg	67.60
			1g	243.00
H30019	2-Aminothiazole-5-carbonitrile, 98% [51640-52-9], C ₃ H ₃ N ₃ S, F.W. 125.20, m.p. ca 185°, UN3439  R:20/21/22, S:36/37		1g	163.00
H51809	2-Aminothiobenzamide, 97% [2454-39-9], C ₇ H ₆ N ₂ S, F.W. 152.21, m.p. 115-120°, MDL MFCD00963496  R:22-36/37/38, S:26-36/37		1g	53.00
			5g	197.00
H51806	3-Aminothiobenzamide, 97% [78950-36-4], C ₇ H ₆ N ₂ S, F.W. 152.21, m.p. 127-131°, MDL MFCD04973325  R:22-36/37/38, S:26-36/37		1g	57.00
			5g	213.00
H51849	4-Aminothiobenzamide, 97% [4114-67-4], C ₇ H ₆ N ₂ S, F.W. 152.21, m.p. 201-204°, MDL MFCD00040927  R:22-36/37/38, S:26-36/37		1g	57.00
			5g	213.00
H32109	3-Amino-2-(trifluoromethyl)pyridine, 97% [106877-32-1], C ₆ H ₅ F ₃ N ₂ , F.W. 162.11, m.p. 68-72°, UN2811, MDL MFCD09753488  R:21-25-36/37/38, S:26-36/37-45		250mg	100.00
			1g	280.00
H30382	Ammonia, 7M in methanol [7864-41-7], NH ₃ , F.W. 17.03, f.p. 14°(57°F), d. 0.779, UN2924,   R:11-23/24/25-34-39/23/24/25, S:9-16-26-36/37/39-45		100ml	67.00
			1L	210.00
H32592	Ammonium O,O'-dimethyldithiophosphate, 95% [1066-97-3], C ₂ H ₁₀ NO ₂ PS ₂ , F.W. 175.21, m.p. 146-150°, EINECS 213-922-5  R:22-36/37/38, S:26-36/37		1g	34.30
			10g	215.00

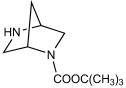
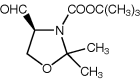
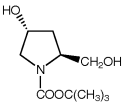
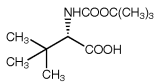
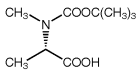
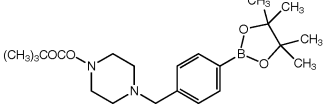
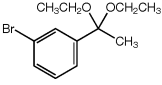
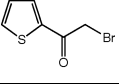
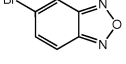
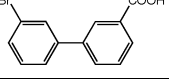
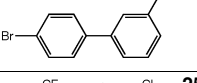
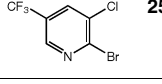
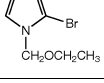
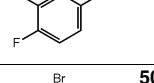
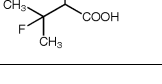
Stock #	Description		Size	\$
H51921	Anthracene-9,10-diboric acid bis(pinacol) ester, 95% [863992-56-7], C ₂₈ H ₃₂ B ₂ O ₄ , F.W. 430.16		1g 5g	48.80 195.00
H31944	Arsenic(V) oxide, 99.6% (metals basis) [1303-28-2], As ₂ O ₅ , F.W. 229.79, m.p. 315° dec., d. 4.32, UN1559, EINECS 215-116-9, RTECS CG2275000, †		25g 100g	89.80 300.00
45616	Arsenic(III) sulfide, 99.9% (metals basis) [1303-33-9], As ₂ S ₃ , F.W. 246.03, m.p. 300°, b.p. 707°, d. 3.43, Merck 14,806, UN1557, EINECS 215-117-4, MDL MFCD00003435, †		5g 25g	29.90 132.00
45522	Arsenic(III) sulfide, 99.999% (metals basis) [Arsenic trisulfide] [1303-33-9], As ₂ S ₃ , F.W. 246.03, m.p. 300°, b.p. 707°, d. 3.43, Merck 14,806, UN1557, EINECS 215-117-4, MDL MFCD00003435, †		5g 25g	63.00 177.00
H51064	1-Aza-15-crown-5, 97%  [66943-05-3], C ₁₀ H ₂₁ NO ₄ , F.W. 219.28, EINECS 266-523-3, BRN 971959, MDL MFCD00075465		250mg 1g 5g	39.00 115.00 467.00
H51065	1-Aza-18-crown-6, 95%  [33941-15-0], C ₁₂ H ₂₅ NO ₅ , F.W. 263.33, m.p. 46-49°, EINECS 251-751-8, BRN 1074079, MDL MFCD00075466		250mg 1g 5g	43.60 127.00 514.00
H30237	7-Azaindole-5-carbonitrile, 97% [5-Cyano-7-azaindole, 1H-Pyrrolo 2,3-b]pyridine-5-carbonitrile] [517918-95-5], C ₈ H ₅ N ₃ , F.W. 143.15, m.p. 226-230°		250mg 1g	74.60 207.00
45552	Barium dodecairon nonadecaoxide [Barium ferrite] [11138-11-7], BaFe ₁₂ O ₁₉ , F.W. 1111.49, m.p. >1315°, UN1564, EINECS 234-387-4, MDL MFCD00075637, †		5g 25g 100g	17.00 60.00 215.00
H51103	Benzaldehyde-d1, 95%  [3592-47-0], C ₆ H ₅ CDO, F.W. 107.13, m.p. -26°, b.p. 178-179°, f.p. 63°(145°F), d. 1.055, n _D ²⁰ 1.5450, UN1990, MDL MFCD00084120		1g 5g	72.00 296.00
H51707	1,4-Benzenediboric acid bis(pinacol) ester, 97% C ₁₈ H ₂₈ B ₂ O ₄ , F.W. 330.03		1g 5g	57.50 230.00
H51837	Benzene-1,2-dithiocarboxamide, 97% C ₆ H ₆ N ₂ S ₂ , F.W. 196.28, UN2811		1g 5g	48.00 179.00
H51853	Benzene-1,3-dithiocarboxamide, 97% C ₆ H ₆ N ₂ S ₂ , F.W. 196.28, UN2811		1g 5g	48.00 179.00
H51857	Benzene-1,4-dithiocarboxamide, 97% [13363-51-4], C ₆ H ₆ N ₂ S ₂ , F.W. 196.28, UN2811		1g 5g	57.00 213.00

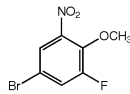
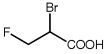
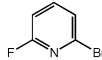
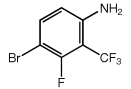
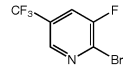
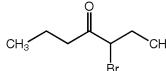
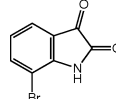
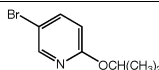
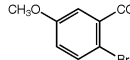
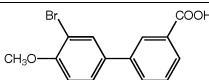
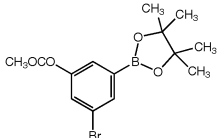
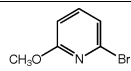
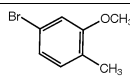
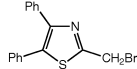
Stock #	Description		Size	\$
H51797	3-(2-Benzimidazolyl)benzamidoxime, 97% C ₁₄ H ₁₂ N ₄ O, F.W. 252.28 ✗ R:36/37/38, S:26-37		250mg 1g	88.40 303.00
H51840	4-(2-Benzimidazolyl)benzamidoxime, 97% C ₁₄ H ₁₂ N ₄ O, F.W. 252.28 ✗ R:36/37/38, S:26-37		250mg 1g	88.00 303.00
H30110	Benzophenone imine, 97% [1013-88-3], C ₁₃ H ₁₁ N, F.W. 181.23, b.p. 151-153°/10mm, d. 1.08, n _D ²⁰ 1.6180 ✗ R:36/37/38, S:26-37		5g 25g	52.40 133.00
H51876	Benzo[b]thiophene-3-boronic acid pinacol ester, 95% [171364-86-6], C ₁₄ H ₁₇ BO ₂ S, F.W. 260.16		250mg 1g	103.00 332.00
H30821	4-Benzylamino-1-butanol, 98+% [59578-63-1], C ₆ H ₅ CH ₂ NH(CH ₂) ₄ OH, F.W. 179.26, b.p. 140°/0.8mm, d. 1.02 ✗ R:36/38, S:26-37		10g 50g	58.10 195.00
H32523	Benzylboronic acid pinacol ester, 96% [87100-28-5], C ₁₃ H ₁₉ BO ₂ , F.W. 218.1, b.p. 65°/0.15mm, d. 0.98		1g 5g 25g	41.10 124.00 470.00
H31126	Benzyl ethyl malonate, 96% [Malonic acid benzyl ethyl ester] [42998-51-6], C ₁₂ H ₁₄ O ₄ , F.W. 222.24		5g 25g	105.00 421.00
H30995	(R)-(-)-1-Benzyl-3-hydroxypiperidine, 97% [(R)-1-Benzyl-3-piperidinol] [91599-81-4], C ₁₂ H ₁₇ NO, F.W. 191.27, MDL MFCD00274307 ✗ R:36/37/38, S:26-37		250mg 1g	55.40 130.00
H30615	(S)-(+)-1-Benzyl-3-hydroxypiperidine, 97% [(S)-1-Benzyl-3-piperidinol] [91599-79-0], C ₁₂ H ₁₇ NO, F.W. 191.27, MDL MFCD03426359 ✗ R:36/37/38, S:26-37		250mg 1g	61.00 154.00
H51675	1-Benzyl-3-iodo-7-azaindole, 97% ▲ C ₁₄ H ₁₁ IN ₂ , F.W. 334.16 ✗ R:36/37/38, S:26-37		5g	180.00
H51154	Benzylmagnesium chloride, 1M in MeTHF ▲ ▽ [6921-34-2], C ₆ H ₅ CH ₂ MgCl, F.W. 150.89, f.p. -17° (1.4°F), UN2924, t ⚠️ R:11-14-19-34, S:8-16-26-36/37/39-43-45		100ml 500ml	33.20 138.00
H30169	1-Benzyl-3-methylpiperazine hydrochloride, 98+% C ₁₂ H ₁₈ N ₂ ·HCl, F.W. 226.75 ✗ R:36/37/38, S:26-37		1g 5g 25g	33.80 112.00 374.00
H30053	4-Benzyloxy-1-bromo-2-fluorobenzene, 98% [185346-79-6], C ₁₃ H ₁₀ BrFO, F.W. 281.13, m.p. 33-35° ✗ R:36/37/38, S:26-37		5g 25g	71.20 237.00
H31488	4-Benzyloxy-2-fluorobenzoic acid, 98% [114045-96-4], C ₁₄ H ₁₁ FO ₃ , F.W. 246.24, m.p. 165-168° ✗ R:36/37/38, S:26-37		1g 5g	44.80 149.00
H51708	4-Benzyloxy-3-formylbenzeneboronic acid pinacol ester ▲ C ₂₀ H ₂₃ BO ₄ , F.W. 338.21		1g 5g	72.50 290.00

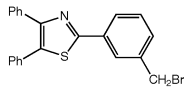
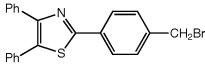
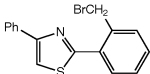
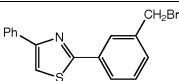
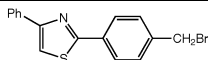
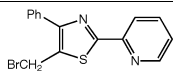
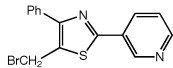
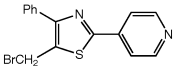
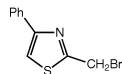
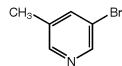
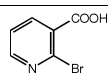
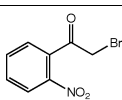
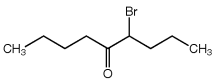
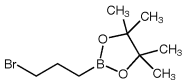
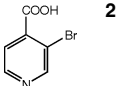
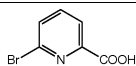
Stock #	Description		Size	\$
H51089	5-Benzyloxytryptamine hydrochloride, 98% [52055-23-9], C ₁₇ H ₁₈ N ₂ O·HCl, F.W. 302.80, EINECS 257-626-4, MDL MFCD00012685		250mg 1g	54.40 133.00
H51844	Benzyl thioacetimidate hydrochloride, 96% [32894-07-8], C ₉ H ₁₁ NS·HCl, F.W. 201.71, MDL MFCD05865210 R:41, S:26-39		1g 5g 25g	17.00 75.00 300.00
H51872	Benzyltriethylammonium dichloroiodate, 97% ▲ (CH ₃ CH ₂) ₃ N(CH ₂ C ₆ H ₅)I ₂ Cl ₂ , F.W. 390.13, m.p. 79-81° R:36/37/38, S:26-37		1g 5g 25g	20.00 75.00 300.00
H31264	(±)-1,1'-Bi(2-naphthylamine), 97% [(±)-1,1'-Binaphthyl-2,2'-diamine, (±)-BINAM] [4488-22-6], C ₂₀ H ₁₆ N ₂ , F.W. 284.36, m.p. 188-193°, RTECS DU3090000, MDL MFCD00145204 R:36/37/38, S:26-37		1g 5g	71.40 235.00
H51087	(±)-1,1'-Bi(2-naphthyl) dimethanesulfonate, 97% △ [182568-57-6], C ₂₂ H ₁₈ O ₂ S ₂ , F.W. 442.50, MDL MFCD05865182		1g 5g	86.60 353.00
H32320	Biphenyl-2-boronic acid pinacol ester, 97% [914675-52-8], C ₁₈ H ₂₁ BO ₂ , F.W. 280.17, m.p. 76-79°		1g 5g 25g	38.70 147.00 588.00
H51761	2-[(4-Biphenyl)dimethylsilyl]benzyl alcohol, 95% C ₂₇ H ₂₂ OSi, F.W. 318.48, Note: Product of Advanced Molecular Technologies Pty Ltd. R:36/37/38, S:26-37		1g 5g	43.00 170.00
H51039	2,5-Bis(bromomethyl)-4-(2-ethylhexyloxy)anisole, 98% [209625-37-6], C ₁₇ H ₂₈ Br ₂ O ₂ , F.W. 422.20, UN3261, MDL MFCD03093944 R:34, S:26-36/37/39-45		1g 5g	114.00 460.00
H51041	2,5-Bis(chloromethyl)-4-(2-ethylhexyloxy)anisole, 98% [146370-52-7], C ₁₇ H ₂₆ Cl ₂ O ₂ , F.W. 333.29, UN1759, MDL MFCD03093939 R:34, S:26-36/37/39-45		1g 5g	113.00 460.00
45523	Bis(1,5-cyclooctadiene)rhodium(I) hexafluoroantimonate [130296-28-5], C ₁₆ H ₂₄ F ₆ RhSb, F.W. 555.01, UN3077, MDL MFCD11113031 R:20/22-51/53, S:61		250mg 1g	69.90 250.00
H51067	2,5-Bis(3,7-dimethyloctyloxy)-terephthalaldehyde, 98% [325461-35-6], C ₂₈ H ₄₀ O ₄ , F.W. 446.66, MDL MFCD03427237 R:36/37/38, S:26-37		250mg 1g	63.70 186.00
H31306	2,2'-Bis(diphenylphosphino)biphenyl, 98% [BIPHEP] [84783-64-2], C ₃₆ H ₂₈ P ₂ , F.W. 522.57, m.p. 211-215° R:36/37/38, S:26-37		250mg 1g 5g	58.50 162.00 540.00
H31441	4,4'-Bis(diphenylphosphinoyl)-2,2',6,6'-tetramethoxy-3,3'-bipyridine, 95% [220998-37-8], C ₃₈ H ₃₄ N ₂ O ₆ P ₂ , F.W. 676.63 R:36/37/38, S:26-37		1g 5g	112.00 448.00

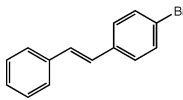
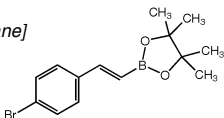
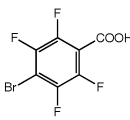
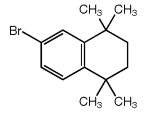
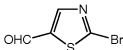
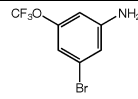
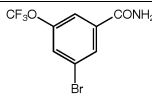
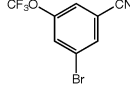
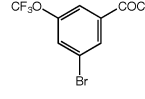
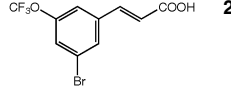
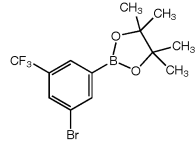
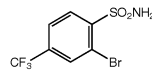
Stock #	Description		Size	\$
H51904	1,4-Bis(2-hydroxy-3,5-di-tert-butylbenzyl)piperazine, 95% [110546-20-8], C ₃₄ H ₅₄ N ₂ O ₂ , F.W. 522.81 ✗ R:36/37/38, S:26-37		1g 5g	72.50 290.00
H51081	4-[Bis(2-hydroxyethyl)amino]benzylidenemalononitrile, 95% [63619-34-1], C ₁₄ H ₁₅ N ₃ O ₂ , F.W. 257.29, MDL MFCD06200699 ✗ R:43, S:24-37		1g 5g	104.00 424.00
H31791	(R)-(+)-2,2'-Bis(methoxymethoxy)-1,1'-binaphthyl, 97% [173831-50-0], C ₂₄ H ₂₂ O ₄ , F.W. 374.44, m.p. 101-105°		250mg 1g	92.10 350.00
H32889	(S)-(-)-2,2'-Bis(methoxymethoxy)-1,1'-binaphthyl, 97% [142128-92-5], C ₂₄ H ₂₂ O ₄ , F.W. 374.44, m.p. 101-105°		250mg 1g	92.10 350.00
H32745	(R)-(+)-2,2'-Bis(methoxymethoxy)-1,1'-binaphthyl-3,3'-diboronic acid pinacol ester, 97% [260442-17-9], C ₃₆ H ₄₄ B ₂ O ₈ , F.W. 626.36, m.p. 193-195°		250mg 1g	150.00 403.00
H32005	(S)-(-)-2,2'-Bis(methoxymethoxy)-1,1'-binaphthyl-3,3'-diboronic acid pinacol ester, 97% [260441-99-4], C ₃₆ H ₄₄ B ₂ O ₈ , F.W. 626.36, m.p. 193-195°		250mg 1g	150.00 403.00
H31081	(1R,2R)-Bis(4-methoxyphenyl)-1,2-ethanediamine, 98% Δ [(1R,2R)-Bis(4-methoxyphenyl)-1,2-diaminoethane, (R,R)-DAEN] [58520-03-9], C ₁₆ H ₂₀ N ₂ O ₂ , F.W. 272.34, m.p. 89-91° ✗ R:36/37/38, S:26-37		1g 5g 25g	59.60 196.00 650.00
H51133	Bis(4-morpholinyl)methane, 98% [5625-90-1], C ₉ H ₁₈ N ₂ O ₂ , F.W. 186.25, b.p. 122-124°/12mm, MDL MFCD00023369 ✗ R:38-41, S:26-37/39		5g 25g	20.10 75.40
45500	Bismuth 2-ethylhexanoate, typically 92%, in 2-ethylhexanoic acid [2-Ethylhexanoic acid bismuth salt] [67874-71-9], C ₂₄ H ₄₅ BiO ₅ , F.W. 638.60, MDL MFCD00043803, † ✗ R:63, S:36/37		5g 25g 100g	14.50 57.80 197.00
45582	Bismuth(III) oxide, NanoArc® BI-7300, 99.5+% [1304-76-3], Bi ₂ O ₃ , F.W. 465.96, m.p. 817°, b.p. ca 1890°, d. 8.9, Merck 14,1273, Application(s): X-ray opaque particle. NanoArc® bismuth oxide can be compounded into polymers or coating systems to increase opacity to x-rays, while the small particle size ensures minimal disruption of physical properties., EINECS 215-134-7, MDL MFCD00003462 Note: Sold in collaboration with Nanophase Technologies Corporation, † ✗ R:36/37/38, S:26-37		25g 100g 500g	24.50 88.00 395.00

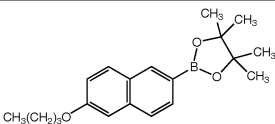
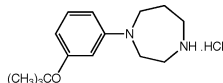
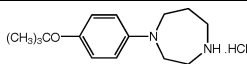
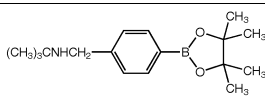
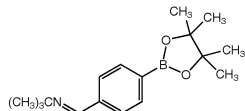
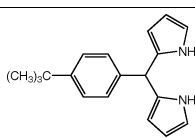
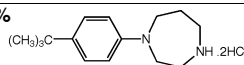
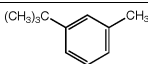
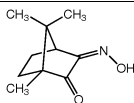
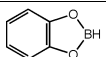
Stock #	Description		Size	\$
H31436	1,4-Bis(trans-2-phenylethenyl)benzene, 97% [1,4-Di(trans- β -styryl)benzene, trans-4- β -Styryl-trans stilbene] [1608-41-9], C ₂₂ H ₁₈ , F.W. 282.39, m.p. 264-266°		1g	196.00
H30338	1,4-Bis(2-phenylethyl)benzene, 97% [1,4-Diphenethylbenzene, 4-Phenethylbibenzyl] [1985-58-6], C ₂₂ H ₂₂ , F.W. 286.42, m.p. 88-90°		1g	176.00
H30395	1,4-Bis(phenylethynyl)benzene, 97% [1849-27-0], C ₂₂ H ₁₄ , F.W. 278.36, m.p. 176-178°		1g 5g 25g	28.60 95.60 319.00
H32985	2,4-Bis(trifluoromethyl)cinnamic acid, JRD, 97% C ₁₅ H ₈ F ₆ O ₂ , F.W. 284.15 ✗ R:36/37/38, S:26-37		250mg 1g	54.40 182.00
H32238	2,5-Bis(trifluoromethyl)cinnamic acid, JRD, 97% C ₁₅ H ₈ F ₆ O ₂ , F.W. 284.15 ✗ R:36/37/38, S:26-37		250mg 1g	54.40 182.00
H32595	3-[2,4-Bis(trifluoromethyl)phenyl]propionic acid, JRD, 97% C ₁₇ H ₈ F ₆ O ₂ , F.W. 286.17 ✗ R:36/37/38, S:26-37		250mg 1g	67.70 225.00
H32656	2-[2,4-Bis(trifluoromethyl)phenyl]thioacetamide, 97% C ₁₀ H ₇ F ₆ NS, F.W. 287.23, m.p. 88-92° ✗ R:22-36/37/38, S:26-36/37		250mg 1g	179.00 495.00
H51709	3-[2-(Boc-amino)acetamido]benzeneboronic acid pinacol ester, 97% C ₁₈ H ₂₉ BN ₂ O ₅ , F.W. 376.26		1g 5g	71.30 280.00
H51816	3-(Boc-amino)benzamidoxime, 97% C ₁₂ H ₁₇ N ₃ O ₃ , F.W. 251.29 ✗ R:36/37/38, S:26-37		250mg 1g	74.00 254.00
H51810	4-(Boc-amino)benzamidoxime, 97% C ₁₂ H ₁₇ N ₃ O ₃ , F.W. 251.29 ✗ R:36/37/38, S:26-37		250mg 1g	74.00 254.00
H51918	4'-(Boc-amino)biphenyl-3-carboxylic acid, 95% C ₁₈ H ₁₉ NO ₄ , F.W. 313.35		1g 5g	58.10 230.00
H51789	6-(Boc-amino)nicotinamidoxime, 97% C ₁₁ H ₁₆ N ₃ O ₃ , F.W. 252.27, m.p. 120-125° ✗ R:36/37/38, S:26-37		250mg 1g	86.70 277.00
H30741	2-(Boc-amino)pyridine, 95% [2-(tert-Butoxycarbonylamino)pyridine] [38427-94-0], C ₁₀ H ₁₄ N ₂ O ₂ , F.W. 194.23, m.p. 92-94° ✗ R:36/37/38, S:26-37		250mg 1g 5g	38.60 107.00 353.00
H30820	4-(Boc-amino)pyridine, 95% [4-(tert-Butoxycarbonylamino)pyridine] [98400-69-2], C ₁₀ H ₁₄ N ₂ O ₂ , F.W. 194.23, m.p. 147-150° ✗ R:36/37/38, S:26-36		5g 25g	67.60 225.00
H51729	(S)-(-)-3-(Boc-amino)pyrrolidine, 98% Δ [122536-76-9], C ₈ H ₁₅ N ₂ O ₂ , F.W. 186.26, m.p. 50-54°, [α] _D ²⁰ -23° (c=1 in methanol), UN3259, BRN 5377813, MDL MFCD00143194 ✗ R:34, S:26-36/37/39-45		5g 25g	79.20 312.00
H51676	9-Boc-7-bromo-6-chloro-7-deazapurine, 97% C ₁₁ H ₁₁ BrClN ₃ O ₂ , F.W. 332.58		1g 5g	88.50 350.00

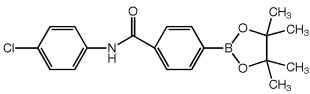
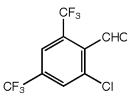
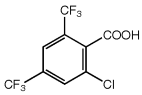
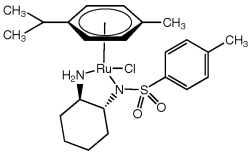
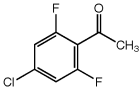
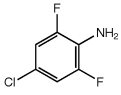
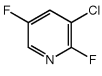
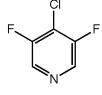
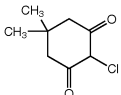
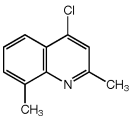
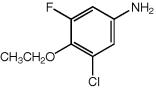
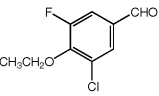
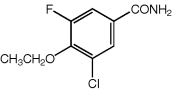
Stock #	Description	Size	\$
H51097	(1S,4S)-(-)-2-Boc-2,5-diazabicyclo[2.2.1]heptane, 98% $\triangle$ [113451-59-5], C ₁₀ H ₁₅ N ₂ O ₂ , F.W. 198.26, m.p. 74-76°, [α] _D ²⁰ -44° (c=1 in chloroform), UN3259, MDL MFCD01569250 	250mg 1g 5g	53.10 114.00 510.00
H51049	(S)-(-)-3-Boc-2,2-dimethyloxazolidine-4-carboxaldehyde, 95% [102308-32-7], C ₁₁ H ₁₉ NO ₄ , F.W. 229.27, BRN 3591324, MDL MFCD00209557 	250mg 1g 5g	52.20 148.00 517.00
H51050	N-Boc-trans-4-hydroxy-L-prolinol, 96% [61478-26-0], C ₁₀ H ₁₉ NO ₄ , F.W. 217.26, MDL MFCD02094386 	1g 5g	109.00 441.00
H51136	N-Boc-L-tert-leucine, 98% [62965-35-9], C ₁₁ H ₂₁ NO ₄ , F.W. 231.29, m.p. 118-121°, MDL MFCD00065574 	1g 5g 25g	21.10 70.50 230.00
H31317	N-Boc-N-methyl-L-alanine, 98% [Boc-N-Me-Ala-OH, N-tert-Butoxycarbonyl-N-methyl-L-alanine] [16948-16-6], C ₉ H ₁₇ NO ₄ , F.W. 203.24, m.p. 88-92°, [α] _D ²⁰ -31° (c=1 in ethanol) 	1g 5g	70.70 236.00
H51743	4-(4-Boc-1-piperazinylmethyl)benzeneboronic acid pinacol ester, 95% C ₂₅ H ₃₅ BN ₂ O ₄ , F.W. 402.34 	1g 5g	78.80 315.00
45573	Boron nitride, low binder, 99.5% (metals basis), Hot pressed rod, 6.4mm dia x 203mm long [10043-11-5], BN, F.W. 24.82, Merck 14,1346, EINECS 233-136-6, MDL MFCD00011317, †	1pc 5pcs	170.00 750.00
45575	Boron nitride, low binder, 99.5% (metals basis), Hot pressed rod, 9.5mm dia x 203mm long [10043-11-5], BN, F.W. 24.82, Merck 14,1346, EINECS 233-136-6, MDL MFCD00011317, †	1pc 5pcs	209.00 940.00
H32157	3'-Bromoacetophenone diethyl ketal, 98% [480439-43-8], C ₁₂ H ₁₇ BrO ₂ , F.W. 273.17 	5g 25g	55.30 185.00
H51060	2-(Bromoacetyl)thiophene, 97% $\blacksquare$ [10531-41-6], C ₆ H ₅ BrOS, F.W. 205.07, m.p. 29-33°, UN3261, MDL MFCD02677721 	1g 5g	174.00 660.00
H51674	5-Bromobenzofurazan, 97% C ₆ H ₃ BrN ₂ O, F.W. 199.01 	1g	40.00
H51766	3'-Bromobiphenyl-3-carboxylic acid, 95% C ₁₃ H ₉ BrO ₂ , F.W. 277.12 	1g 5g	60.80 195.00
H51762	4'-Bromobiphenyl-3-carboxylic acid, 95% C ₁₃ H ₉ BrO ₂ , F.W. 277.12 	1g 5g	60.80 195.00
H32055	2-Bromo-3-chloro-5-(trifluoromethyl)pyridine, 97% [75806-84-7], C ₆ H ₂ BrClF ₃ N, F.W. 260.44, b.p. 80°/32mm, MDL MFCD00153072 	250mg 1g 5g	99.20 207.00 425.00
H51860	2-Bromo-1-(ethoxymethyl)imidazole [850429-54-8], C ₆ H ₈ BrN ₂ O, F.W. 205.05 	1g 5g	146.00 563.00
H30604	3-Bromo-4-fluorobenzyl alcohol, 97% [77771-03-0], C ₇ H ₆ BrFO, F.W. 205.03, b.p. 80°/0.5mm 	5g 25g	59.30 197.00
H32246	2-Bromo-3-fluoroisovaleric acid, 97% C ₅ H ₈ BrFO ₂ , F.W. 199.02, m.p. 52-55°, UN3261, MDL MFCD08704576 	500mg 2g	72.10 203.00

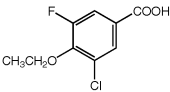
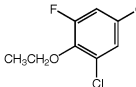
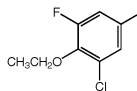
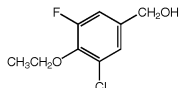
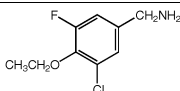
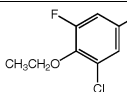
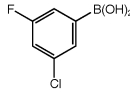
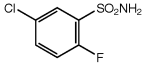
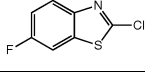
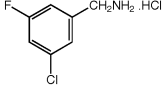
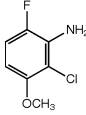
Stock #	Description		Size	\$
H30220	4-Bromo-2-fluoro-6-nitroanisole, 97% [74266-66-3], C ₆ H ₃ BrFNO ₂ , F.W. 250.02, m.p. 53-56° ✗ R:36/37/38, S:26-37		5g	155.00
			25g	619.00
H31718	2-Bromo-3-fluoropropionic acid, 96% C ₃ H ₄ BrFO ₂ , F.W. 170.97, b.p. 62-63°/0.2mm, UN3265, MDL MFCD09800640 ✗ R:34, S:26-36/37/39-45		250mg	140.00
			1g	392.00
H31981	2-Bromo-3-fluoropyridine, 97% [40273-45-8], C ₅ H ₃ BrFN, F.W. 175.99, UN2810 ✗ R:20/21/22-36/37/38, S:26-36/37		1g	86.10
			5g	287.00
H30924	2-Bromo-6-fluoropyridine, 97% [144100-07-2], C ₅ H ₃ BrFN, F.W. 175.99, m.p. 30-34°, UN2811 ✗ R:20/21/22-36/37/38, S:26-36/37		1g	79.50
			5g	264.00
H31969	4-Bromo-3-fluoro-2-(trifluoromethyl)aniline, JRD, 97% C ₇ H ₄ BrFN, F.W. 258.01, UN2810 ✗ R:20/21/22-36/38, S:26-36/37		250mg	95.20
			1g	317.00
H31993	2-Bromo-3-fluoro-5-(trifluoromethyl)pyridine, 97% C ₆ H ₃ BrF ₂ N, F.W. 243.99 ✗ R:36/38, S:26-37		250mg	75.60
			1g	210.00
H32094	3-Bromo-4-heptanone, 98% [42330-10-9], C ₇ H ₁₃ BrO, F.W. 193.08, b.p. 39°/1mm, f.p. 75°(167°F), d. 1.248, n _D ²⁰ 1.4580, UN3265 ✗ R:34, S:26-36/37/39-45		1g	59.70
			10g	373.00
H32843	7-Bromoisatin, 97% [20780-74-9], C ₈ H ₅ BrNO ₂ , F.W. 226.03, m.p. 195-199° ✗ R:36/37/38, S:26-37		1g	62.80
			10g	395.00
H32564	5-Bromo-2-isopropoxy pyridine, 97% [870521-31-6], C ₈ H ₁₀ BrNO, F.W. 216.08, n _D ²⁰ 1.5305 ✗ R:36/38, S:26-37		1g	97.00
			5g	323.00
H31030	Bromomalonaldehyde, 97% Δ [2065-75-0], BrCH(CHO) ₂ , F.W. 150.96, m.p. ca 145° dec., EINECS 430-470-6 ✗ R:22-41, S:26-39		1g	31.00
			5g	102.00
			50g	367.00
H31250	2-Bromo-5-methoxybenzamide, 98% [99848-43-8], C ₈ H ₈ BrNO ₂ , F.W. 230.66, m.p. 159-161° ✗ R:36/37/38, S:26-37		5g	54.10
			25g	180.00
H51912	3'-Bromo-4'-methoxybiphenyl-3-carboxylic acid, 95% C ₁₄ H ₁₁ BrO ₃ , F.W. 307.14 ✗ R:36/37/38, S:26-37		1g	48.80
			5g	195.00
H51718	3-Bromo-5-(methoxycarbonyl)benzeneboronic acid pinacol ester, 97% [1025718-78-8], C ₁₄ H ₁₈ BBrO ₄ , F.W. 341.01, m.p. 98-102°		250mg	45.00
			1g	155.00
H30593	2-Bromo-6-methoxypyridine [40473-07-2], C ₆ H ₆ BrNO, F.W. 188.02, b.p. 206°, f.p. 104°(219°F), d. 1.530, n _D ²⁰ 1.5600, MDL MFCD00088345 ✗ R:36/37/38, S:26-37		5g	31.10
			25g	107.00
H30391	5-Bromo-2-methylanisole, 97% [4-Bromo-2-methoxytoluene] [67868-73-9], C ₈ H ₇ BrO, F.W. 201.06, b.p. 102-104°/13mm, n _D ²⁰ 1.5630		1g	35.40
			5g	117.00
H51812	2-Bromomethyl-4,5-diphenylthiazole, 97% C ₁₈ H ₁₂ BrNS, F.W. 330.24 ✗ R:36/37/38, S:26-37		250mg	33.40
			1g	115.00

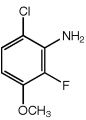
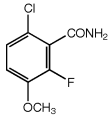
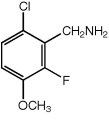
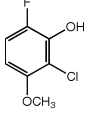
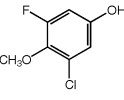
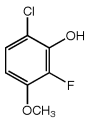
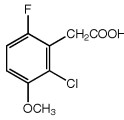
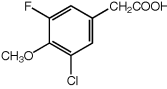
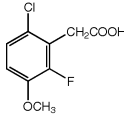
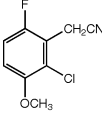
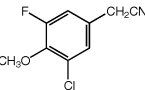
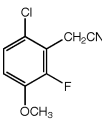
Stock #	Description		Size	\$
H51800	2-[3-(Bromomethyl)phenyl]-4,5-diphenylthiazole, 97% C ₂₂ H ₁₆ BrNS, F.W. 406.34 R:36/37/38, S:26-37		250mg 1g	44.20 152.00
H51841	2-[4-(Bromomethyl)phenyl]-4,5-diphenylthiazole, 97% C ₂₂ H ₁₆ BrNS, F.W. 406.34, UN3261 R:34, S:26-36/37/39-45		250mg 1g	44.00 152.00
H51832	2-[2-(Bromomethyl)phenyl]-4-phenylthiazole, 97% C ₁₆ H ₁₂ BrNS, F.W. 330.24, UN3261 R:34, S:26-36/37/39-45		1g 5g	79.00 294.00
H51834	2-[3-(Bromomethyl)phenyl]-4-phenylthiazole, 97% C ₁₆ H ₁₂ BrNS, F.W. 330.24, UN3261 R:34, S:26-36/37/39-45		1g 5g	79.00 294.00
H51836	2-[4-(Bromomethyl)phenyl]-4-phenylthiazole, 97% C ₁₆ H ₁₂ BrNS, F.W. 330.24, UN3261 R:34, S:26-36/37/39-45		1g 5g	79.00 294.00
H51772	5-Bromomethyl-4-phenyl-2-(2-pyridyl)thiazole, 97% C ₁₅ H ₁₁ BrN ₂ S, F.W. 331.23, UN3261 R:34, S:26-36/37/39-45		1g 5g	102.00 380.00
H51779	5-Bromomethyl-4-phenyl-2-(3-pyridyl)thiazole, 97% C ₁₅ H ₁₁ BrN ₂ S, F.W. 331.23, UN3261 R:34, S:26-36/37/39-45		1g 5g	99.40 370.00
H51791	5-Bromomethyl-4-phenyl-2-(4-pyridyl)thiazole, 97% C ₁₅ H ₁₁ BrN ₂ S, F.W. 331.23, UN3261 R:34, S:26-36/37/39-45		1g 5g	99.40 370.00
H51780	2-Bromomethyl-4-phenylthiazole, 97% [78502-79-1], C ₁₀ H ₈ BrNS, F.W. 254.14, UN3261 R:34, S:26-36/37/39-45		1g 5g	63.40 236.00
H30137	3-Bromo-5-methylpyridine, 97% [5-Bromo-3-picoline] [3430-16-8], C ₆ H ₆ BrN, F.W. 172.03, d. 1.49 R:20/21/22-36/37/38, S:26-36/37		1g 5g	110.00 364.00
H30239	2-Bromonicotinic acid, 97% [3-Bromopyridine-3-carboxylic acid] [35905-85-2], C ₆ H ₄ BrNO ₂ , F.W. 202.01, m.p. ca 254° R:36/37/38, S:26-37		1g 5g	38.00 141.00
H30275	2-Bromo-2'-nitroacetophenone, 98% [2-Nitrophenacyl bromide] [6851-99-6], C ₈ H ₆ BrNO ₂ , F.W. 244.05, m.p. 53-57°, UN3261 R:34, S:26-36/37/39-45		1g 5g	31.40 96.00
H32136	4-Bromo-5-nonanone, 98% [42330-11-0], C ₉ H ₁₇ BrO, F.W. 221.14, b.p. 62°/1mm, f.p. 92°(198°F), d. 1.174, n _D ²⁰ 1.4600, UN3265 R:34, S:26-36/37/39-45		1g 10g	87.70 548.00
H30646	3-Bromopropylboronic acid pinacol ester, 98% [2-(3-Bromopropyl)-4,4,5,5-tetramethyl-1,3,2-dioxaborolane] [124215-44-7], C ₉ H ₁₈ BBrO ₂ , F.W. 248.96, b.p. 61-64°/0.35mm, d. 1.177 R:36/38, S:26-37		1g 5g 25g	31.90 128.00 510.00
H31047	3-Bromopyridine-4-carboxylic acid, 97% [3-Bromoisonicotinic acid] [13959-02-9], C ₆ H ₄ BrNO ₂ , F.W. 202.01, m.p. ca 235° dec. R:36/37/38, S:26-37		250mg 1g	70.70 196.00
H30553	6-Bromopyridine-2-carboxylic acid, 97% [6-Bromopicolinic acid] [21190-87-4], C ₆ H ₄ BrNO ₂ , F.W. 202.01, m.p. 188-191° R:36/37/38, S:26-37		1g 5g	44.60 172.00

Stock #	Description		Size	\$
H30189	4-Bromo-trans-stilbene, 98% [(E)-1-Bromo-4-(2-phenylethenyl)benzene] [13041-70-8], C ₁₄ H ₁₁ Br, F.W. 259.15, m.p. 138-140° ☒ R:36/37/38, S:26-37		1g 5g	79.40 241.00
H31412	trans-4-Bromo-β-styrylboronic acid pinacol ester, 96% [2-[(E)-2-(4-Bromophenyl)vinyl]-1,3,2-dioxaborolane] C ₁₄ H ₁₈ BBro ₂ , F.W. 309.01, m.p. 85-87° ☒ R:36/37/38, S:26-37		250mg 1g 5g	86.70 241.00 726.00
H32982	4-Bromo-2,3,5,6-tetrafluorobenzoic acid, 98% [4707-24-8], C ₇ HBrF ₄ O ₂ , F.W. 272.98, m.p. 142-145° ☒ R:36/37/38, S:26-37		1g 5g	39.60 131.00
H30436	6-Bromo-1,1,4,4-tetramethyl-1,2,3,4-tetrahydronaphthalene, 98% [27452-17-1], C ₁₄ H ₁₉ Br, F.W. 267.21, m.p. 34-37°		1g 10g	53.00 393.00
H31096	2-Bromothiazole-5-carboxaldehyde, 95% △ [464192-28-7], C ₄ H ₂ BrNOS, F.W. 192.03, m.p. 108-110° ☒ R:36/37/38, S:26-37		250mg 1g	55.50 154.00
H32510	3-Bromo-5-(trifluoromethoxy)aniline, JRD, 97% C ₈ H ₆ BrF ₃ NO, F.W. 256.02, UN2810 ☒ R:20/21/22-36/38, S:26-28-36/37		1g 5g	130.00 520.00
H32805	3-Bromo-5-(trifluoromethoxy)benzamide, JRD, 97% C ₉ H ₆ BrF ₃ NO ₂ , F.W. 284.03 ☒ R:36/37/38, S:26-37		1g 5g	110.00 450.00
H31778	3-Bromo-5-(trifluoromethoxy)benzonitrile, JRD, 97% C ₈ H ₅ BrF ₃ NO, F.W. 266.01, m.p. 35-37°, UN3439 ☒ R:20/21/22-36/38, S:26-36/37		1g 5g	130.00 520.00
H31775	3-Bromo-5-(trifluoromethoxy)benzoyl chloride, JRD, 97% ▢ C ₈ H ₅ BrClF ₃ O ₂ , F.W. 303.46, UN3265 ☒ R:34, S:26-36/37/39-45		1g 5g	98.50 395.00
H32952	3-Bromo-5-(trifluoromethoxy)cinnamic acid, JRD, 97% C ₁₀ H ₆ BrF ₃ O ₃ , F.W. 311.05 ☒ R:36/37/38, S:26-37		250mg 1g	54.40 180.00
H51713	3-Bromo-5-(trifluoromethyl)benzeneboronic acid pinacol ester, 97% [479411-92-2], C ₁₃ H ₁₃ BBrF ₃ O ₂ , F.W. 350.97, m.p. 50-52°		250mg 1g	60.00 190.00
H31649	2-Bromo-4-(trifluoromethyl)benzenesulfonamide, 97% [351003-63-9], C ₇ H ₅ BrF ₃ NO ₂ S, F.W. 304.08, m.p. 160-164°, MDL MFCD03094287 ☒ R:36/37/38, S:26-37		1g 5g	90.50 359.00
H51869	2-Bromo-1-tritylimidazole, 97% [67478-47-1], C ₂₂ H ₁₇ BrN ₂ , F.W. 389.29, m.p. 204-206° ☒ R:36/37/38, S:26-37		1g 5g	130.00 500.00
H51171	2-Butenylmagnesium chloride, 0.5M in MeTHF △ ▢ [22649-70-3], CH ₃ CH=CHCH ₂ MgCl, F.W. 114.86, UN3399 ☒ R:11-14/15-19-34, S:8-16-26-36/37/39-43-45		100ml 500ml	107.00 400.00

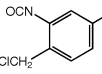
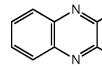
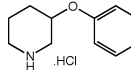
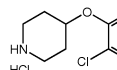
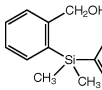
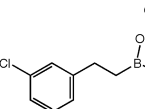
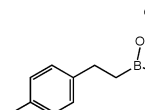
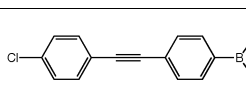
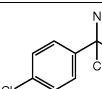
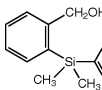
Stock #	Description		Size	\$
H51693	6-n-Butoxynaphthalene-2-boronic acid pinacol ester, 97% C ₂₀ H ₂₇ BO ₃ , F.W. 326.24		1g 5g	58.50 230.00
H51752	1-(3-tert-Butoxyphenyl)homopiperazine monohydrochloride, 98% [934992-04-8], C ₁₅ H ₂₄ N ₂ O·HCl, F.W. 284.83, m.p. 143-147°, Note: Sold in collaboration with Tosoh Organic Chemical Co., Ltd.		500mg	390.00
H51687	1-(4-tert-Butoxyphenyl)homopiperazine monohydrochloride, 98% [934991-96-5], C ₁₅ H ₂₄ N ₂ O·HCl, F.W. 284.83, m.p. 197-201°, Note: Sold in collaboration with Tosoh Organic Chemical Co., Ltd.		500mg	390.00
H51669	4-(tert-Butylaminomethyl)benzeneboronic acid pinacol ester C ₁₇ H ₂₈ BNO ₂ , F.W. 289.22		1g 5g	40.50 156.00
H51122	4-(tert-Butyliminomethyl)benzeneboronic acid pinacol ester C ₁₇ H ₂₈ BNO ₂ , F.W. 287.20, MDL MFCD11113039		1g 5g 25g	28.40 109.00 439.00
H51164	tert-Butylmagnesium chloride, 1M in MeTHF   [677-22-5], (CH ₃) ₃ CMgCl, F.W. 116.87, UN3399		100ml 500ml	36.00 135.00
H51066	α-(4-tert-Butylphenyl)di(2-pyrrolyl)methane, 99%  [167482-98-6], C ₁₉ H ₂₂ N ₂ , F.W. 278.39, m.p. 158-162°, MDL MFCD03454220   R:36/37/38, S:26-37		1g 5g	111.00 446.00
H51702	1-(4-tert-Butylphenyl)homopiperazine dihydrochloride, 98% [934992-03-7], C ₁₅ H ₂₄ N ₂ ·2HCl, F.W. 305.30, m.p. 240-244°, Note: Sold in collaboration with Tosoh Organic Chemical Co., Ltd.   R:36/37/38, S:26-37		500mg	390.00
H51048	tert-Butyl propiolate, 98% [13831-03-3], HC≡CCO ₂ C(CH ₃) ₃ , F.W. 126.15, UN3272, BRN 1747175, MDL MFCD00060100   R:10-36/37/38, S:26-37		1g 5g	108.00 462.00
H32668	3-tert-Butyltoluene, 99% [1075-38-3], C ₁₁ H ₁₆ , F.W. 148.24, m.p. -41°, b.p. 189°, d. 0.87, UN2667   R:20/22-36/37-51/53, S:26-36-61		1g 5g	134.00 466.00
H30770	tert-Butyl vinyl ether, 98%, stab. with ca 0.1% N,N-diethylaniline [2-Methyl-2-vinyloxypropane] [926-02-3], (CH ₃) ₃ COCH=CH ₂ , F.W. 100.16, b.p. 75-76°, f.p. -17°(1°F), d. 0.762, n _D ²⁰ 1.3990, UN3271  R:11-52/53, S:9-16-33-61		5g 25g	51.70 157.00
H51034	(1R,E)-(+)-Camphorquinone 3-oxime, 99% [31571-14-9], C ₁₀ H ₁₅ NO ₂ , F.W. 181.23, BRN 3200377, MDL MFCD00074848		1g 5g	141.00 569.00
45606	m-Carborane, 98% [1,7-Dicarbadoecaborane, m-Barene] [16986-24-6], C ₂ H ₁₂ B ₁₀ , F.W. 144.25, MDL MFCD00151574,  R:22-36/37/38, S:26-36/37		500mg 2g 10g	48.60 152.00 608.00
H30234	Catecholborane, 50% w/w in toluene (99+% dry wt.) [1,3,2-Benzodioxaborole] [274-07-7], C ₆ H ₅ BO ₂ , F.W. 119.92, UN2924, EINECS 205-991-5,    R:11-14-34-48/20-63-67, S:16-26-30-36/37/39-45		25g 100g	132.00 361.00
H31733	Celatom® FW-14 [68855-54-9], EINECS 272-489-0, MDL MFCD00132803   R:40-48/20, S:22-36		250g 1kg	20.80 60.50
45657	Cerium(IV) isopropoxide isopropanol adduct, Ce 31.5-32.4%  Ce(OC ₃ H ₇) ₄ ·C ₃ H ₇ OH, F.W. 436.57, UN3180   R:11-34, S:16-26-36/37/39-45		1g 5g	59.00 235.00

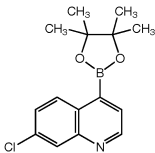
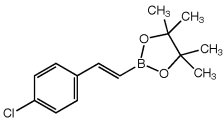
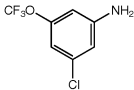
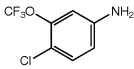
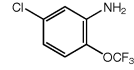
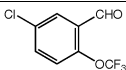
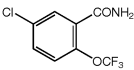
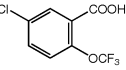
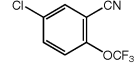
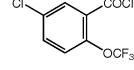
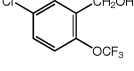
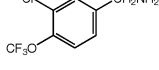
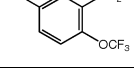
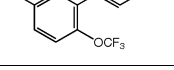
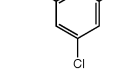
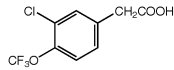
Stock #	Description		Size	\$
H51919	4'-Chlorobenzanilide-4-boronic acid pinacol ester, 95% C ₁₅ H ₂₁ BClNO ₃ , F.W. 357.64		1g 5g	48.80 195.00
H31532	2-Chloro-4,6-bis(trifluoromethyl)benzaldehyde, JRD, 97%  C ₉ H ₅ ClF ₆ O, F.W. 276.56  R:36/38, S:26-37		250mg 1g	95.20 317.00
H31784	2-Chloro-4,6-bis(trifluoromethyl)benzoic acid, JRD, 97% C ₉ H ₅ ClF ₆ O ₂ , F.W. 292.56  R:36/37/38, S:26-37		250mg 1g	95.20 317.00
H30264	Chloro(p-cymene)-N-(p-toluenesulfonyl)-(R,R)-1,2-cyclohexanediamineruthenium(I), 97% [(S,S)-TsDACH RuCl(p-cymene)], (p-Cymene)N-(p-toluenesulfonyl)-(R,R)-1,2-cyclohexanediamineruthenium(I) chloride [213603-12-4], C ₂₅ H ₃₃ ClN ₂ O ₂ SuRuS, F.W. 538.11		250mg 1g	114.00 344.00
H32711	4'-Chloro-2',6'-difluoroacetophenone, JRD, 97% [1017777-45-5], C ₈ H ₅ ClF ₂ O, F.W. 190.57  R:36/38, S:26-37		1g 5g	105.00 415.00
H32371	4-Chloro-2,6-difluoroaniline, JRD, 97% C ₆ H ₄ ClF ₂ N, F.W. 163.55, m.p. 47-50°, UN2811  R:20/21/22-36/38, S:26-36/37		250mg 1g	54.40 182.00
H32640	3-Chloro-2,5-difluoropyridine, 97% [851178-97-7], C ₅ H ₂ ClF ₂ N, F.W. 149.53, n _D ²⁰ 1.480, UN1992  R:10-20/21/22-36/38, S:26-36/37		250mg 1g	66.70 185.00
H31611	4-Chloro-3,5-difluoropyridine, 97% [851178-97-7], C ₅ H ₂ ClF ₂ N, F.W. 149.53, n _D ²⁰ 1.4795, UN1992  R:10-20/21/22-36/38, S:26-36/37		250mg 1g	60.70 179.00
H51035	2-Chloro-5,5-dimethyl-1,3-cyclohexanedione, 98% [7298-89-7], C ₈ H ₁₁ ClO ₂ , F.W. 174.62, BRN 2326749, MDL MFCD00010502		250mg 1g 5g	52.20 134.00 545.00
H51714	4-Chloro-2,8-dimethylquinoline [32314-39-9], C ₁₁ H ₁₀ ClN, F.W. 191.66, MDL MFCD00272394  R:36/37/38, S:26-37		250mg 1g	73.20 255.00
H32000	3-Chloro-4-ethoxy-5-fluoroaniline, JRD, 97% C ₉ H ₉ ClFNO, F.W. 189.61, UN2811  R:20/21/22-33-36/38, S:26-36/37		250mg 1g	54.40 182.00
H32761	3-Chloro-4-ethoxy-5-fluorobenzaldehyde, JRD, 97%  C ₉ H ₈ ClFO ₂ , F.W. 202.61  R:36/37/38, S:26-37		1g 5g	25.60 105.00
H32959	3-Chloro-4-ethoxy-5-fluorobenzamide, JRD, 97% C ₉ H ₉ ClFNO ₂ , F.W. 217.62  R:36/37/38, S:26-37		1g 5g	81.60 325.00

Stock #	Description		Size	\$
H32037	3-Chloro-4-ethoxy-5-fluorobenzoic acid, JRD, 97% <chem>C9H8ClFO3</chem> , F.W. 218.61 ✗ R:36/37/38, S:26-37		1g	25.60
			5g	102.00
H31890	3-Chloro-4-ethoxy-5-fluorobenzonitrile, JRD, 97% <chem>C9H7ClFNO</chem> , F.W. 199.61, UN3439 ✗ R:20/21/22-36/37/38, S:26-36/37		1g	99.20
			5g	397.00
H32411	3-Chloro-4-ethoxy-5-fluorobenzoyl chloride, JRD, 97% ▢ <chem>C9H7Cl2FO2</chem> , F.W. 237.06, UN3265 ✗ R:34, S:26-36/37/39-45		1g	78.10
			5g	315.00
H31990	3-Chloro-4-ethoxy-5-fluorobenzyl alcohol, JRD, 97% <chem>C9H10ClFO2</chem> , F.W. 204.63 ✗ R:36/38, S:26-37		1g	81.60
			5g	326.00
H31681	3-Chloro-4-ethoxy-5-fluorobenzylamine, JRD, 97% △ <chem>C9H11ClFNO</chem> , F.W. 203.64, UN2735 ✗ R:34, S:26-36/37/39-45		250mg	67.70
			1g	230.00
H32653	3-Chloro-4-ethoxy-5-fluorophenol, JRD, 97% <chem>C9H8ClFO2</chem> , F.W. 190.60 ✗ R:22-36/37/38, S:26-36/37		250mg	54.40
			1g	182.00
H51870	2-Chloro-1-(ethoxymethyl)imidazole [850429-55-9], <chem>C6H9ClN2O</chem> , F.W. 160.60 ✗ R:36/38, S:26-37		250mg	51.00
			1g	150.00
H51144	3-Chloro-2-fluoroaniline, 97% [2106-04-9], <chem>C6H5ClFN</chem> , F.W. 145.56, b.p. 214°, f.p. >110°(230°F), d. 1.324, n _D ²⁰ 1.564, UN2810, MDL MFCD00069415, † ✗ R:20/21/22-33-36/38, S:26-28-36/37		5g	14.10
			25g	45.00
			100g	150.00
H32137	3-Chloro-5-fluorobenzeneboronic acid, 95% [328956-61-2], <chem>C6H5BClFO2</chem> , F.W. 174.37, MDL MFCD06801741 ✗ R:36/37/38, S:26-37		1g	126.00
			5g	420.00
H32144	5-Chloro-2-fluorobenzenesulfonamide, 97% [351003-57-1], <chem>C6H5ClFNO2S</chem> , F.W. 209.62, m.p. 134-136°, MDL MFCD03094289 ✗ R:36/37/38, S:26-37		1g	84.50
			5g	280.00
H31443	3-Chloro-2-fluorobenzonitrile, 97% [94087-40-8], <chem>C7H3ClFN</chem> , F.W. 155.56, m.p. 40-44°, f.p. 73°(163°F), UN3439, EINECS 301-934-4 ✗ R:20/21/22-36/38, S:26-36/37		1g	42.00
			5g	168.00
H32383	2-Chloro-6-fluorobenzothiazole, 98% [399-74-6], <chem>C7H3ClFNS</chem> , F.W. 187.62, m.p. 89-93° ✗ R:36/37/38, S:26-37		1g	67.20
			5g	225.00
H51131	3-Chloro-5-fluorobenzylamine hydrochloride <chem>C7H8ClFN.HCl</chem> , F.W. 196.05, MDL MFCD11113045 ✗ R:36/37/38, S:26-37		1g	105.00
			5g	424.00
H32742	2-Chloro-6-fluoro-3-methoxyaniline, JRD, 97% <chem>C7H7ClFNO</chem> , F.W. 175.59, UN2811 ✗ R:20/21/22-33-36/38, S:26-36/37		1g	117.00
			5g	465.00

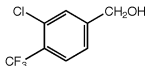
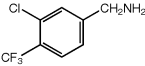
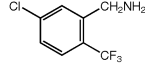
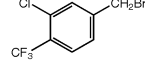
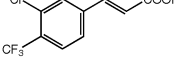
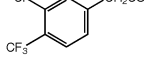
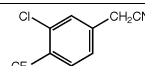
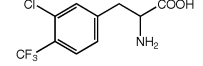
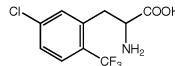
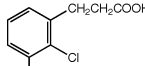
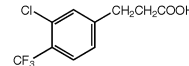
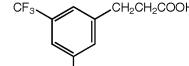
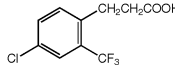
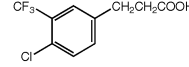
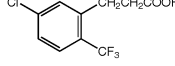
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H31779	6-Chloro-2-fluoro-3-methoxyaniline, JRD, 97% [1017777-77-3], C ₇ H ₆ ClFNO, F.W. 175.59, m.p. 40-43°, UN2811  R:20/21/22-33-36/38, S:26-28-36/37		1g	117.00
			5g	470.00
H31638	6-Chloro-2-fluoro-3-methoxybenzamide, JRD, 97% [886761-75-7], C ₈ H ₇ ClFNO ₂ , F.W. 203.60  R:36/37/38, S:26-37		1g	25.60
			5g	102.00
H31546	6-Chloro-2-fluoro-3-methoxybenzylamine, JRD, 97%  C ₉ H ₉ ClFNO, F.W. 189.61, m.p. 55-57°, UN3259  R:34, S:26-36/37/39-45		1g	88.60
			5g	355.00
H32204	2-Chloro-6-fluoro-3-methoxyphenol, JRD, 97% C ₇ H ₆ ClFO ₂ , F.W. 176.57, m.p. 66-68°  R:21/22-36/37/38, S:26-36/37		1g	117.00
			5g	467.00
H31801	3-Chloro-5-fluoro-4-methoxyphenol, JRD, 97% C ₇ H ₆ ClFO ₂ , F.W. 176.57, m.p. 62-65°  R:21/22-36/37/38, S:26-36/37		250mg	54.40
			1g	182.00
H32654	6-Chloro-2-fluoro-3-methoxyphenol, JRD, 97% C ₇ H ₆ ClFO ₂ , F.W. 176.57, m.p. 56-59°  R:21/22-36/37/38, S:26-36/37		1g	117.00
			5g	470.00
H31665	2-Chloro-6-fluoro-3-methoxyphenylacetic acid, JRD, 97% C ₉ H ₇ ClFO ₃ , F.W. 218.61  R:36/37/38, S:26-37		250mg	67.70
			1g	226.00
H32062	3-Chloro-5-fluoro-4-methoxyphenylacetic acid, JRD, 97% C ₉ H ₇ ClFO ₃ , F.W. 218.61  R:36/37/38, S:26-37		250mg	81.70
			1g	273.00
H32712	6-Chloro-2-fluoro-3-methoxyphenylacetic acid, JRD, 97% C ₉ H ₇ ClFO ₃ , F.W. 218.61  R:36/37/38, S:26-37		250mg	67.70
			1g	230.00
H31966	2-Chloro-6-fluoro-3-methoxyphenylacetonitrile, JRD, 97% C ₉ H ₇ ClFNO, F.W. 199.61, m.p. 53-55°, UN3439  R:20/21/22-36/37/38, S:26-36/37		250mg	54.40
			1g	182.00
H31666	3-Chloro-5-fluoro-4-methoxyphenylacetonitrile, JRD, 97% C ₉ H ₇ ClFNO, F.W. 199.61, UN3439  R:20/21/22-36/37/38, S:26-36/37		250mg	67.70
			1g	225.00
H32287	6-Chloro-2-fluoro-3-methoxyphenylacetonitrile, JRD, 97% C ₉ H ₇ ClFNO, F.W. 199.61, m.p. 85-87°, UN3439  R:20/21/22-36/37/38, S:26-36/37		250mg	54.40
			1g	182.00

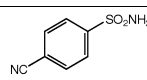
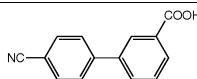
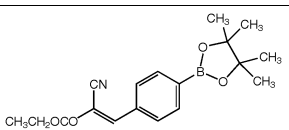
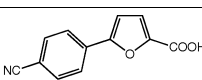
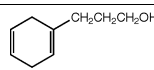
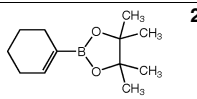
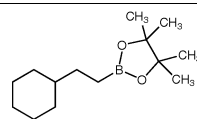
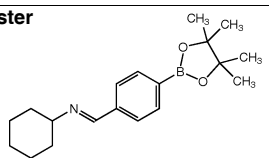
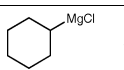
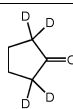
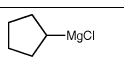
Stock #	Description		Size	\$
H51879	2-Chloro-4-fluoro-3-methylbenzoic acid, 97% C ₈ H ₆ ClFO ₂ , F.W. 188.58, m.p. 186-189° ✗ R:36/37/38, S:26-37		250mg 1g	95.00 306.00
H30410	3-Chloro-5-fluorophenylacetonitrile, 97% [493038-93-0], C ₈ H ₅ ClFN, F.W. 169.59, UN3276 ✗ R:20/21/22-36/38, S:26-36/37		1g 5g	87.80 292.00
H31742	2-(3-Chloro-2-fluorophenyl)ethanol, JRD, 94% C ₈ H ₈ ClFO, F.W. 174.60, n _D ²⁰ 1.5305		250mg 1g	33.60 112.00
H32291	2-(3-Chloro-4-fluorophenyl)ethanol, JRD, 96% C ₈ H ₈ ClFO, F.W. 174.60		250mg 1g	33.60 112.00
H30170	3-Chloro-5-fluoropyridine, 98% [514797-99-0], C ₅ H ₃ ClFN, F.W. 131.54, m.p. 25-27°, UN2811 ✗ R:20/21/22-36/37/38, S:26-36/37		1g 5g	141.00 470.00
H32281	3-Chloro-2-fluoro-5-(trifluoromethyl)aniline, JRD, 97% C ₇ H ₄ ClF ₃ N, F.W. 213.56, UN2810 ✗ R:20/21/22-36/38, S:26-28-36/37		250mg 1g	47.60 159.00
H32861	3-Chloro-2-fluoro-6-(trifluoromethyl)aniline, JRD, 97% C ₇ H ₄ ClF ₃ N, F.W. 213.56, UN2810 ✗ R:20/21/22-36/38, S:26-28-36/37		250mg 1g	47.60 160.00
H32693	3-Chloro-2-fluoro-6-(trifluoromethyl)benzonitrile, JRD, 97% C ₈ H ₃ ClF ₃ N, F.W. 223.55, UN3439 ✗ R:20/21/22-36/37/38, S:26-36/37		250mg 1g	47.60 160.00
H51863	2-Chloroimidazole, 97% [16265-04-6], C ₃ H ₃ ClN ₂ , F.W. 102.52, m.p. 165-166°, MDL MFCD02179530 ✗ R:22-37/38-41, S:26-36/37/39		250mg 1g	67.00 196.00
H51128	6-Chloro-7-iodo-7-deazapurine ▲ [123148-78-7], C ₆ H ₃ ClIN ₃ , F.W. 279.47, m.p. ca 180° dec., MDL MFCD09263258 ✗ R:36/37/38, S:26-37		1g 5g	93.40 358.00
H31342	3-Chloro-4-methylaniline hydrochloride, 97% [7745-89-3], C ₇ H ₈ ClN·HCl, F.W. 178.06, m.p. ca 260° subl., EINECS 202-446-3, RTECS XU5425000 ✗ R:20/21/22-33-36/37/38, S:26-36/37		100g 500g	27.70 92.20
H30893	3-Chloro-2-methylbenzeneboronic acid, 97% [3-Chloro-2-methylphenylboronic acid] [313545-20-9], C ₇ H ₆ BClO ₂ , F.W. 170.40 ✗ R:36/37/38, S:26-37		250mg 1g	129.00 359.00
H51867	2-Chloro-1-methylimidazole [253453-91-7], C ₄ H ₅ ClN ₂ , F.W. 116.55, MDL MFCD02179531 ✗ R:36/37/38, S:26-37		250mg 1g	51.00 150.00
H30636	2-Chloromethyl-3-methyl-4-(2,2,2-trifluoroethoxy)pyridine hydrochloride, 97% [127337-60-4], C ₉ H ₉ ClF ₃ NO·HCl, F.W. 276.09, m.p. 208-212° ✗ R:36/37/38, S:26-37		1g	132.00
H31108	6-Chloro-2-methyl-3-nitropyridine, 98% [6-Chloro-5-nitro-2-picoline] [22280-60-0], C ₆ H ₅ ClN ₂ O ₂ , F.W. 172.57, m.p. 54-58° ✗ R:22-36/37/38, S:26-36/37		5g	187.00

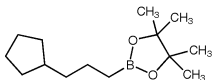
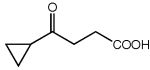
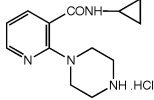
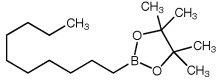
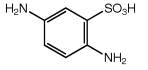
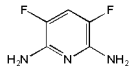
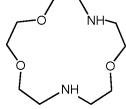
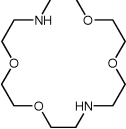
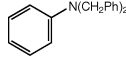
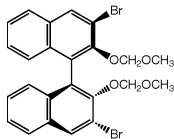
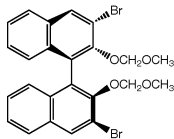
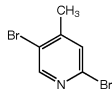
Stock #	Description		Size	\$
H51056	4-Chloromethyl-1,3-phenylene diisocyanate, 97%  [51979-57-8], C ₆ H ₃ ClN ₂ O ₂ , F.W. 208.60, UN2922, MDL MFCD00134404   R:20/22-34-42, S:23-26-36/37/39-45		250mg 1g 5g	44.10 111.00 448.00
H51099	4-Chloro-3-methylphenyl isocyanate, 98%  [51488-20-1], C ₈ H ₆ ClNO, F.W. 167.60, n _D ²⁰ 1.5580, UN2206  R:23-36/37/38-42, S:23-26-36/37-45		1g 10g	37.80 235.00
H51734	2-Chloro-3-methylquinoxaline, 95% [32601-86-8], C ₉ H ₇ ClN ₂ , F.W. 178.62, m.p. 71-75°  R:36/37/38, S:26-37		250mg 1g	72.00 207.00
H51180	3-(3-Chlorophenoxy)piperidine hydrochloride C ₁₁ H ₁₄ ClNO·HCl, F.W. 248.15, MDL MFCD11870724  R:36/37/38, S:26-37		250mg 1g	62.60 226.00
H51029	4-(2-Chlorophenoxy)piperidine hydrochloride [849107-20-6], C ₁₁ H ₁₄ ClNO·HCl, F.W. 248.15, MDL MFCD02684107  R:36/37/38, S:26-37		250mg 1g	59.90 216.00
H51058	4-Chlorophenylacetylene, 98% [873-73-4], C ₈ H ₅ Cl, F.W. 136.58, UN1325, MDL MFCD00191917   R:11-36/37/38, S:26-37		1g 5g	132.00 536.00
H51736	2-[(4-Chlorophenyl)dimethylsilyl]benzyl alcohol, 95% C ₁₅ H ₁₇ ClOSi, F.W. 276.83, Note: Product of Advanced Molecular Technologies Pty Ltd.  R:36/37/38, S:26-37		1g 5g	54.40 220.00
H30464	2-(3-Chlorophenyl)ethylboronic acid pinacol ester, 97% [2-[2-(3-Chlorophenyl)ethyl]-4,4,5,5-tetramethyl-1,3,2-dioxaborolane] C ₁₄ H ₂₀ BClO ₂ , F.W. 266.58, n _D ²⁰ 1.4965		1g 5g	47.60 153.00
H30303	2-(4-Chlorophenyl)ethylboronic acid pinacol ester, 98% [2-[2-(4-Chlorophenyl)ethyl]-4,4,5,5-tetramethyl-1,3,2-dioxaborolane] [444094-88-6], C ₁₄ H ₂₀ BClO ₂ , F.W. 266.58, m.p. 34-38°		1g 5g	44.60 149.00
H51915	4-(4-Chlorophenylethynyl)benzeneboronic acid pinacol ester, 95% C ₂₀ H ₂₀ BClO ₂ , F.W. 338.64		1g 5g	48.80 195.00
H51161	3-Chlorophenylmagnesium bromide, 1M in MeTHF  [36229-42-2], C ₆ H ₄ BrClMg, F.W. 215.76, UN2924   R:11-14-19-34, S:8-16-26-36/37/39-43-45		50ml 100ml	112.00 179.00
H51163	4-Chlorophenylmagnesium bromide, 1M in MeTHF  [873-77-8], C ₆ H ₄ BrClMg, F.W. 215.76, UN2924   R:11-14-19-34, S:8-16-26-36/37/39-43-45		100ml 500ml	34.50 129.00
H32527	1-(4-Chlorophenyl)-1-methylethylamine, 97%  [17797-11-4], C ₉ H ₁₂ ClN, F.W. 169.66, b.p. 43-45°/0.3mm, n _D ²⁰ 1.5330, UN2735  R:34, S:26-36/37/39-45		250mg 1g	95.70 266.00
H32301	4-Chloropyridine-2-carboxylic acid, 94% [4-Chloropicolinic acid] [5470-22-4], C ₆ H ₄ ClNO ₂ , F.W. 157.56, m.p. ca 180° dec.  R:36/37/38, S:26-37		1g 10g	43.80 273.00
H30923	5-Chloropyridine-2-carboxylic acid, 95% [5-Chloropicolinic acid] [86873-60-1], C ₆ H ₄ ClNO ₂ , F.W. 157.56, m.p. 166-171°  R:36/37/38, S:26-37		1g 10g	69.00 431.00
H51738	2-(6-Chloro-3-pyridyl)dimethylsilyl]benzyl alcohol, 95% C ₁₄ H ₁₆ ClNOSi, F.W. 277.82, Note: Product of Advanced Molecular Technologies Pty Ltd.  R:36/37/38, S:26-37		1g 5g	67.50 270.00

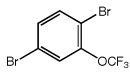
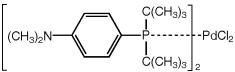
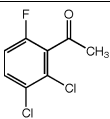
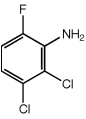
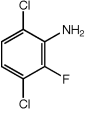
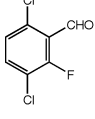
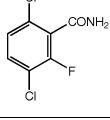
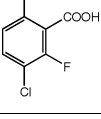
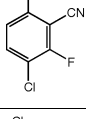
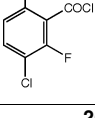
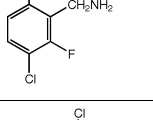
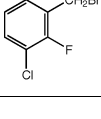
Stock #	Description		Size	\$
H51037	7-Chloroquinoline-4-boronic acid pinacol ester, 96% [871125-83-6], C ₁₅ H ₁₇ BClO ₂ , F.W. 289.56, MDL MFCD07783020 ✗ R:36/37/38, S:26-37		1g	121.00
			5g	484.00
H51692	4-Chloro-β-styrylboronic acid pinacol ester, 97% C ₁₄ H ₁₈ BClO ₂ , F.W. 264.56		1g	60.00
			5g	240.00
H30241	3-Chloro-5-(trifluoromethoxy)aniline, JRD, 97% [151276-13-0], C ₇ H ₅ ClF ₃ NO, F.W. 211.57, UN2810 ✗ R:20/21/22-36/38, S:26-36/37		250mg	55.50
			1g	186.00
H31752	4-Chloro-3-(trifluoromethoxy)aniline, JRD, 97% C ₇ H ₅ ClF ₃ NO, F.W. 211.57, UN2810 ✗ R:20/21/22-36/38, S:26-28-36/37		250mg	67.70
			1g	230.00
H32259	5-Chloro-2-(trifluoromethoxy)aniline, JRD, 97% C ₇ H ₅ ClF ₃ NO, F.W. 211.57, UN2810 ✗ R:20/21/22-36/38, S:26-28-36/37		250mg	81.70
			1g	273.00
H31584	5-Chloro-2-(trifluoromethoxy)benzaldehyde, JRD, 97% △ C ₈ H ₅ ClF ₃ O ₂ , F.W. 224.56 ✗ R:36/38, S:26-37		1g	115.00
			5g	460.00
H32184	5-Chloro-2-(trifluoromethoxy)benzamide, JRD, 97% C ₈ H ₅ ClF ₃ NO ₂ , F.W. 239.58 ✗ R:36/37/38, S:26-37		250mg	67.70
			1g	226.00
H32451	5-Chloro-2-(trifluoromethoxy)benzoic acid, JRD, 97% C ₈ H ₅ ClF ₃ O ₃ , F.W. 240.56 ✗ R:36/37/38, S:26-37		1g	115.00
			5g	460.00
H31977	5-Chloro-2-(trifluoromethoxy)benzonitrile, JRD, 97% C ₈ H ₅ ClF ₃ NO, F.W. 221.56, UN3276 ✗ R:20/21/22-36/38, S:26-36/37		250mg	67.70
			1g	230.00
H32638	5-Chloro-2-(trifluoromethoxy)benzoyl chloride, JRD, 97% ▢ C ₈ H ₅ ClF ₃ O ₂ , F.W. 259.01, UN3265 ☞ R:34, S:26-36/37/39-45		250mg	54.40
			1g	182.00
H32806	5-Chloro-2-(trifluoromethoxy)benzyl alcohol, JRD, 97% C ₈ H ₇ ClF ₃ O ₂ , F.W. 226.58 ✗ R:36/37/38, S:26-37		250mg	54.40
			1g	182.00
H31785	3-Chloro-4-(trifluoromethoxy)benzylamine, JRD, 97% △ C ₈ H ₇ ClF ₃ NO, F.W. 225.60, UN2735 ☞ R:34, S:26-36/37/39-45		250mg	54.40
			1g	182.00
H32620	5-Chloro-2-(trifluoromethoxy)benzyl bromide, JRD, 97% C ₈ H ₇ BrClF ₃ O, F.W. 289.48, UN3265 ☞ R:34, S:26-36/37/39-45		250mg	67.70
			1g	226.00
H31717	5-Chloro-2-(trifluoromethoxy)cinnamic acid, JRD, 97% C ₁₀ H ₆ ClF ₃ O ₃ , F.W. 266.60 ✗ R:36/37/38, S:26-37		250mg	54.40
			1g	182.00
H32579	3-Chloro-5-(trifluoromethoxy)phenol, JRD, 97% C ₇ H ₄ ClF ₃ O ₂ , F.W. 212.55 ✗ R:21/22-36/37/38, S:26-36/37		250mg	33.60
			1g	112.00
H32437	3-Chloro-4-(trifluoromethoxy)phenylacetic acid, JRD, 97% C ₉ H ₆ ClF ₃ O ₃ , F.W. 254.59 ✗ R:36/37/38, S:26-37		250mg	81.70
			1g	275.00

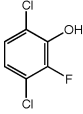
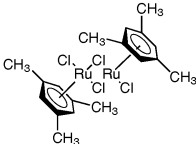
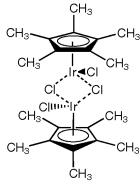
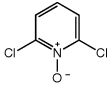
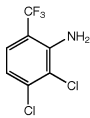
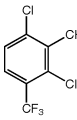
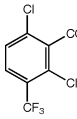
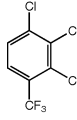
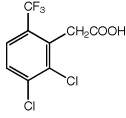
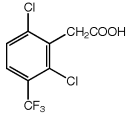
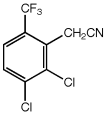
Stock #	Description		Size	\$
H32477	5-Chloro-2-(trifluoromethoxy)phenylacetic acid, JRD, 97% C ₉ H ₆ ClF ₃ O ₃ , F.W. 254.59 ☒ R:36/37/38, S:26-37		250mg 1g	109.00 365.00
H32661	3-Chloro-4-(trifluoromethoxy)phenylacetonitrile, JRD, 97% C ₉ H ₆ ClF ₃ NO, F.W. 235.59, UN3276 ☒ R:20/21/22-36/38, S:26-36/37		250mg 1g	67.70 225.00
H32598	5-Chloro-2-(trifluoromethoxy)phenylacetonitrile, JRD, 97% C ₉ H ₆ ClF ₃ NO, F.W. 235.59, m.p. 64-66°, UN3439 ☒ R:20/21/22-36/37/38, S:26-36/37		250mg 1g	95.20 317.00
H31935	3-[4-Chloro-3-(trifluoromethoxy)phenyl]propionic acid, JRD, 97% C ₁₀ H ₈ ClF ₃ O ₃ , F.W. 268.62 ☒ R:36/37/38, S:26-37		250mg 1g	67.70 226.00
H31610	3-[5-Chloro-2-(trifluoromethoxy)phenyl]propionic acid, JRD, 97% C ₁₀ H ₈ ClF ₃ O ₃ , F.W. 268.62 ☒ R:36/37/38, S:26-37		250mg 1g	67.70 226.00
H32247	2'-Chloro-3'-(trifluoromethyl)acetophenone, JRD, 97% C ₉ H ₆ ClF ₃ O, F.W. 222.59 ☒ R:36/37/38, S:23-26-37		1g 5g	45.30 182.00
H31635	3-Chloro-4-(trifluoromethyl)benzaldehyde, JRD, 97% △ C ₈ H ₄ ClF ₃ O, F.W. 208.56 ☒ R:36/38, S:26-37		1g 5g	45.70 182.00
H32515	2-Chloro-6-(trifluoromethyl)benzamide, JRD, 97% C ₈ H ₅ ClF ₃ NO, F.W. 223.58 ☒ R:36/37/38, S:26-37		1g 5g	78.10 312.00
H31799	3-Chloro-4-(trifluoromethyl)benzamide, JRD, 97% C ₈ H ₅ ClF ₃ NO, F.W. 223.58 ☒ R:36/37/38, S:26-37		1g 5g	106.00 425.00
H51712	3-Chloro-5-(trifluoromethyl)benzeneboronic acid pinacol ester, 97% [942069-65-0], C ₁₃ H ₁₅ BClF ₃ O ₂ , F.W. 306.52		250mg 1g	68.90 220.00
H32637	2-Chloro-4-(trifluoromethyl)benzenesulfonamide, 97% [146533-47-3], C ₇ H ₅ ClF ₃ NO ₂ S, F.W. 259.63, m.p. 168-171°, MDL MFCD00179384 ☒ R:36/37/38, S:26-37		250mg 1g	46.40 130.00
H31789	2-Chloro-4-(trifluoromethyl)benzoic acid, JRD, 97% C ₈ H ₄ ClF ₃ O ₂ , F.W. 224.56 ☒ R:36/37/38, S:26-37		250mg 1g	43.60 145.00
H30490	2-Chloro-6-(trifluoromethyl)benzoic acid, JRD, 97% [6-Chloro-α,α,α-trifluoro-o-toluic acid] [2376-00-3], C ₈ H ₄ ClF ₃ O ₂ , F.W. 224.56, m.p. 123-125° ☒ R:36/37/38, S:26-37		1g 5g	51.40 205.00
H32038	3-Chloro-4-(trifluoromethyl)benzoic acid, JRD, 97% [115754-20-6], C ₈ H ₄ ClF ₃ O ₂ , F.W. 224.56 ☒ R:36/37/38, S:26-37		1g 5g	46.70 187.00
H31731	2-Chloro-6-(trifluoromethyl)benzonitrile, JRD, 97% C ₈ H ₄ ClF ₃ N, F.W. 205.56, UN3439 ☒ R:20/21/22-36/37/38, S:26-36/37		1g 5g	124.00 495.00
H30062	2-Chloro-6-(trifluoromethyl)benzoyl chloride, JRD, 97% ☒ [916420-44-5], C ₈ H ₃ ClF ₃ O, F.W. 243.01, UN3261 ☒ R:34, S:26-36/37/39-45		1g 5g	63.90 255.00

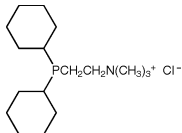
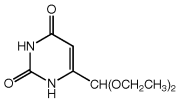
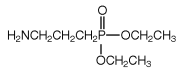
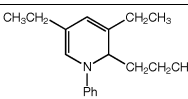
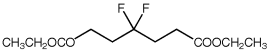
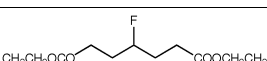
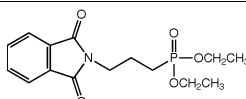
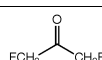
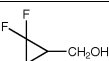
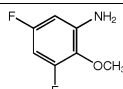
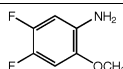
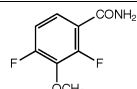
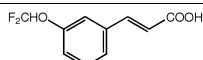
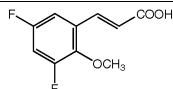
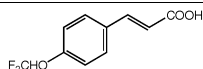
Stock #	Description		Size	\$
H31522	3-Chloro-4-(trifluoromethyl)benzyl alcohol, JRD, 97% C ₈ H ₆ ClF ₃ O, F.W. 210.58  R:36/37/38, S:26-37		1g 5g	114.00 457.00
H31883	3-Chloro-4-(trifluoromethyl)benzylamine, JRD, 97% Δ C ₈ H ₇ ClF ₃ N, F.W. 209.60, UN2735  R:34, S:26-36/37/39-45		1g 5g	122.00 490.00
H32593	5-Chloro-2-(trifluoromethyl)benzylamine, JRD, 97% Δ C ₈ H ₇ ClF ₃ N, F.W. 209.60, UN2735  R:34, S:26-36/37/39-45		1g 5g	110.00 439.00
H32212	3-Chloro-4-(trifluoromethyl)benzyl bromide, JRD, 97% C ₈ H ₆ BrClF ₃ , F.W. 273.48, UN3265  R:34, S:26-36/37/39-45		1g 5g	122.00 488.00
H31572	3-Chloro-4-(trifluoromethyl)cinnamic acid, JRD, 97% C ₁₀ H ₆ ClF ₃ O ₂ , F.W. 250.60, m.p. 183-185°  R:36/37/38, S:26-37		250mg 1g	33.60 112.00
H32819	3-Chloro-4-(trifluoromethyl)phenylacetic acid, JRD, 97% C ₉ H ₆ ClF ₃ O ₂ , F.W. 238.559  R:36/37/38, S:26-37		250mg 1g	45.90 153.00
H30659	2-Chloro-3-(trifluoromethyl)phenylacetonitrile, 97% [22902-81-4], C ₉ H ₆ ClF ₃ N, F.W. 219.59, m.p. 56-59°, UN3439  R:20/21/22-36/38, S:26-36/37		1g 5g	112.00 450.00
H32116	3-Chloro-4-(trifluoromethyl)phenylacetonitrile, JRD, 97% C ₉ H ₆ ClF ₃ N, F.W. 219.59, UN3276  R:20/21/22-36/38, S:26-36/37		250mg 1g	40.80 136.00
H31919	3-Chloro-4-(trifluoromethyl)-DL-phenylalanine, JRD, 97% C ₁₀ H ₉ ClF ₃ NO ₂ , F.W. 267.63		250mg 1g	67.70 226.00
H31947	5-Chloro-2-(trifluoromethyl)-DL-phenylalanine, JRD, 97% C ₁₀ H ₉ ClF ₃ NO ₂ , F.W. 267.63		250mg 1g	67.70 226.00
H32476	3-[2-Chloro-3-(trifluoromethyl)phenyl]propionic acid, JRD, 97% C ₁₀ H ₈ ClF ₃ O ₂ , F.W. 252.62  R:36/37/38, S:26-37		250mg 1g	54.40 182.00
H32245	3-[3-Chloro-4-(trifluoromethyl)phenyl]propionic acid, JRD, 97% C ₁₀ H ₈ ClF ₃ O ₂ , F.W. 252.62  R:36/37/38, S:26-37		250mg 1g	67.70 226.00
H32615	3-[3-Chloro-5-(trifluoromethyl)phenyl]propionic acid, JRD, 97% C ₁₀ H ₈ ClF ₃ O ₂ , F.W. 252.62  R:36/37/38, S:26-37		250mg 1g	67.70 230.00
H32443	3-[4-Chloro-2-(trifluoromethyl)phenyl]propionic acid, JRD, 97% C ₁₀ H ₈ ClF ₃ O ₂ , F.W. 252.62  R:36/37/38, S:26-37		250mg 1g	54.40 182.00
H32460	3-[4-Chloro-3-(trifluoromethyl)phenyl]propionic acid, JRD, 97% C ₁₀ H ₈ ClF ₃ O ₂ , F.W. 252.62  R:36/37/38, S:26-37		250mg 1g	54.40 185.00
H31746	3-[5-Chloro-2-(trifluoromethyl)phenyl]propionic acid, JRD, 97% C ₁₀ H ₈ ClF ₃ O ₂ , F.W. 252.62  R:36/37/38, S:26-37		250mg 1g	67.70 226.00
H51861	2-Chloro-1-tritylimidazole, 97% [67478-48-2], C ₂₂ H ₁₇ ClN ₂ , F.W. 344.84, m.p. 200-202°  R:36/37/38, S:26-37		250mg 1g	43.00 125.00

Stock #	Description	Size	\$	
45579	Cobalt oxide, typically 3.4-4.5%, Molybdenum oxide typically 11.5-14.5% on alumina †	25g	24.80	
		100g	80.80	
		500g	280.00	
		2.5kg	565.00	
45604	Copper germanide, 99.99% (metals basis) CuGe, F.W. 136.16, †	10g	136.00	
		50g	590.00	
45605	Copper germanium selenide, 99.99% (metals basis) CuGeSe, F.W. 215.12, UN3283, † 	10g	157.00	
		50g	705.00	
45660	Copper(I) selenide, 99.5% (metals basis) [20405-64-5], Cu ₂ Se, F.W. 206.05, m.p. 1113°, d. 6.84, Merck 14,2666, UN3077, EINECS 243-796-7, MDL MFCD00049908, † 	10g	69.20	
		50g	244.00	
H31836	4-Cyanobenzenesulfonamide, 97% [3118-68-1], C ₇ H ₆ N ₂ O ₂ S, F.W. 182.20, m.p. 168-171°, MDL MFCD02089633 		1g	126.00
			5g	420.00
H51906	4'-Cyanobiphenyl-3-carboxylic acid, 95% C ₁₆ H ₉ NO ₂ , F.W. 223.23, MDL MFCD04117352 		1g	51.60
			5g	210.00
H51130	4-(2-Cyano-3-ethoxy-3-oxo-1-propen-1-yl)-benzeneboronic acid pinacol ester C ₁₈ H ₂₂ BNO ₄ , F.W. 327.18, MDL MFCD11113044 		1g	25.90
			5g	98.90
			25g	401.00
H31065	4-Cyano-2-iodopyridine, 95% ▲ [2-Iodoisonicotinonitrile, 2-Iodopyridine-4-carbonitrile] [114821-24-8], C ₆ H ₄ IN ₂ , F.W. 230.01, m.p. 77-80° 		1g	148.00
			5g	493.00
H51763	5-(4-Cyanophenyl)-2-furoic acid, 95% C ₁₂ H ₇ NO ₃ , F.W. 213.19 		1g	58.10
			5g	230.00
H31465	3-(1,4-Cyclohexadien-1-yl)-1-propanol, 97%, stab. with 0.1% BHT [3-(3-Hydroxypropyl)-1,4-cyclohexadiene] [87151-66-4], C ₉ H ₁₄ O, F.W. 138.21		1g	66.70
			5g	271.00
H51036	1-Cyclohexene-1-boronic acid pinacol ester, 95% [141091-37-4], C ₁₂ H ₂₁ BO ₂ , F.W. 208.10, MDL MFCD05663845		250mg	70.00
			1g	205.00
H31156	2-Cyclohexylethylboronic acid pinacol ester, 96% [2-(2-Cyclohexylethyl)-4,4,5,5-tetramethyl-1,3,2-dioxaborolane] [167692-95-7], C ₁₄ H ₂₇ BO ₂ , F.W. 238.18		1g	60.00
			5g	200.00
H51127	4-(Cyclohexyliminomethyl)benzeneboronic acid pinacol ester C ₁₉ H ₂₈ BNO ₂ , F.W. 313.24, MDL MFCD11113042		1g	28.40
			5g	109.00
			25g	439.00
H51165	Cyclohexylmagnesium chloride, 1M in MeTHF ▲ ▢ [931-51-1], C ₆ H ₁₁ ClMg, F.W. 142.91, UN2924, † 		100ml	30.00
			500ml	113.00
H51095	Cyclopentanone-2,2,5,5-d₄, 95% [3997-89-5], C ₅ H ₄ D ₄ O, F.W. 88.14, UN2245, MDL MFCD00190415 		1g	213.00
			5g	861.00
H51166	Cyclopentylmagnesium chloride, 1M in MeTHF ▲ ▢ [32916-51-1], C ₅ H ₉ ClMg, F.W. 128.88, UN2924 		100ml	49.30
			500ml	185.00

Stock #	Description		Size	\$
H31158	3-Cyclopentylpropylboronic acid pinacol ester, 97% [2-(3-Cyclopentylpropyl)-4,4,5,5-tetramethyl[1,3,2]dioxaborolane] C ₁₄ H ₂₇ BO ₂ , F.W. 238.18, b.p. 114°/5mm		1g	63.40
			5g	201.00
			25g	678.00
H51158	Cyclopropylmagnesium bromide, 0.5M in MeTHF △ ▢ [23719-80-4], C ₃ H ₅ BrMg, F.W. 145.28, UN3399  R:11-14/15-19-34, S:8-16-26-36/37/39-43-45		50ml	129.00
			100ml	206.00
H51078	4-Cyclopropyl-4-oxobutyric acid, 95% [53712-75-7], C ₇ H ₁₀ O ₃ , F.W. 142.15, MDL MFCD01320151  R:36/37/38, S:26-37		1g	143.00
H51683	N-Cyclopropyl-2-(1-piperazinyl)nicotinamide hydrochloride, 97% C ₁₃ H ₁₈ N ₄ O·HCl, F.W. 282.77		250mg	62.60
			1g	226.00
H30403	1-Decylboronic acid pinacol ester, 98% [2-(n-Decyl)-4,4,5,5-tetramethyl-1,3,2-dioxaborolane] [141091-38-5], C ₁₆ H ₃₃ BO ₂ , F.W. 268.25		1g	39.40
			5g	157.00
			25g	626.00
H31999	2,5-Diaminobenzenesulfonic acid, 97% [88-45-9], C ₆ H ₆ N ₂ O ₃ S, F.W. 188.20, m.p. 298-300°, EINECS 201-832-9, RTECS DB5900100, BRN 2970398, †  R:36/37/38, S:26-37		50g	37.30
			250g	138.00
H30365	2,6-Diamino-3,5-difluoropyridine, 98% [3,5-Difluoro-2,6-pyridinediamine] [247069-27-8], C ₅ H ₅ F ₂ N ₃ , F.W. 145.11, m.p. 158-162°, UN2811  R:24/25-36/37/38, S:26-36/37-45		1g	38.00
			5g	126.00
H51062	1,7-Diaza-15-crown-5, 97% △ [31249-95-3], C ₁₀ H ₂₂ N ₂ O ₅ , F.W. 218.29, EINECS 250-530-3, BRN 607391, MDL MFCD00005109  R:36/37/38, S:26-37		250mg	43.80
			1g	132.00
			5g	540.00
H51063	1,10-Diaza-18-crown-6, 96% △ [23978-55-4], C ₁₂ H ₂₆ N ₂ O ₆ , F.W. 262.35, EINECS 245-965-0, RTECS HM1650000, BRN 609764, MDL MFCD00005112  R:36/37/38, S:26-37		250mg	43.80
			1g	132.00
			5g	540.00
H32091	N,N-Dibenzylaniline, 99% [91-73-6], C ₂₀ H ₁₉ N, F.W. 273.37, m.p. 69-71°, b.p. 180°/1mm, EINECS 202-093-5, MDL MFCD00022015, †  R:21/22, S:36/37		25g	37.80
			100g	105.00
H31557	(R)-3,3'-Dibromo-2,2'-bis(methoxymethoxy)-1,1'-binaphthyl, 97% [211734-49-5], C ₂₄ H ₂₀ Br ₂ O ₄ , F.W. 532.23		250mg	150.00
			1g	403.00
H31783	(S)-3,3'-Dibromo-2,2'-bis(methoxymethoxy)-1,1'-binaphthyl, 97% [142010-87-5], C ₂₄ H ₂₀ Br ₂ O ₄ , F.W. 532.23		250mg	150.00
			1g	403.00
H30457	2,5-Dibromo-4-methylpyridine, 97% [2,5-Dibromo-4-picoline] [3430-26-0], C ₆ H ₅ Br ₂ N, F.W. 250.92, m.p. 36-40°  R:36/37/38, S:26-37		1g	65.50
			5g	208.00
H31708	2,3-Dibromopyridine-4-carboxylic acid, 97% [1020056-98-7], C ₆ H ₃ Br ₂ NO ₂ , F.W. 280.90  R:36/37/38, S:26-37		250mg	101.00
			1g	280.00

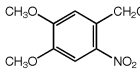
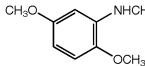
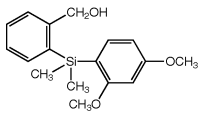
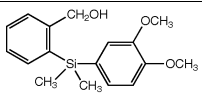
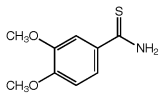
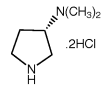
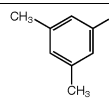
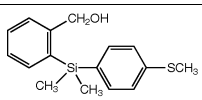
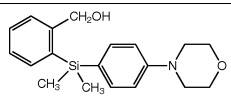
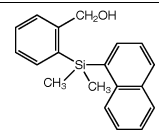
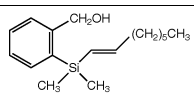
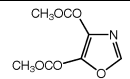
Stock #	Description		Size	\$
H32896	1,4-Dibromo-2-(trifluoromethoxy)benzene, JRD, 97% C ₇ H ₃ Br ₂ F ₃ O, F.W. 319.90 ☒ R:36/37/38, S:23-26-37		1g	28.00
			5g	112.00
45511	Dichlorobis[di-tert-butyl(4-dimethylaminophenyl)-phosphino]palladium(II) [887919-35-9], C ₃₂ H ₅₆ Cl ₂ N ₂ P ₂ Pd, F.W. 708.07 ☒ R:36/37/38, S:26-37		1g	135.00
H32663	2',3'-Dichloro-6'-fluoroacetophenone, JRD, 97% C ₈ H ₅ Cl ₂ FO, F.W. 207.03 ☒ R:36/37/38, S:23-26-37		1g	25.60
			5g	105.00
H31673	2,3-Dichloro-6-fluoroaniline, JRD, 97% C ₆ H ₃ Cl ₂ FN, F.W. 180.01, UN2811 ☒ R:20/21/22-33-36/38, S:26-28-36/37		1g	78.10
			5g	310.00
H32333	3,6-Dichloro-2-fluoroaniline, JRD, 97% C ₆ H ₃ Cl ₂ FN, F.W. 180.01, UN2811 ☒ R:20/21/22-33-36/38, S:26-36/37		1g	85.10
			5g	340.00
H32455	3,6-Dichloro-2-fluorobenzaldehyde, JRD, 97% △ C ₇ H ₃ Cl ₂ FO, F.W. 193.00 ☒ R:36/37/38, S:26-37		1g	39.70
			5g	160.00
H32277	3,6-Dichloro-2-fluorobenzamide, JRD, 97% C ₇ H ₃ Cl ₂ FNO, F.W. 208.02 ☒ R:36/37/38, S:26-37		1g	56.40
			5g	226.00
H32666	3,6-Dichloro-2-fluorobenzoic acid, JRD, 97% C ₇ H ₃ Cl ₂ FO ₂ , F.W. 209.00 ☒ R:36/37/38, S:26-37		1g	39.70
			5g	159.00
H32752	3,6-Dichloro-2-fluorobenzonitrile, JRD, 97% C ₇ H ₂ Cl ₂ FN, F.W. 190.00, UN3439 ☒ R:20/21/22-36/38, S:26-36/37		1g	68.10
			5g	273.00
H32887	3,6-Dichloro-2-fluorobenzoyl chloride, JRD, 97% ▢ C ₇ H ₂ Cl ₂ FO, F.W. 227.45, UN3265 ☒ R:34, S:26-36/37/39-45		1g	45.30
			5g	182.00
H32132	3,6-Dichloro-2-fluorobenzylamine, JRD, 97% △ C ₈ H ₆ Cl ₂ FN, F.W. 194.03, m.p. 44-47°, UN3259 ☒ R:34, S:26-36/37/39-45		250mg	54.40
			1g	182.00
H32932	3,6-Dichloro-2-fluorobenzyl bromide, JRD, 97% C ₈ H ₆ BrCl ₂ F, F.W. 257.92, UN3265 ☒ R:34, S:26-36/37/39-45		1g	78.10
			5g	312.00

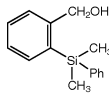
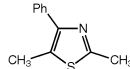
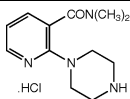
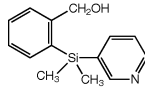
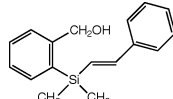
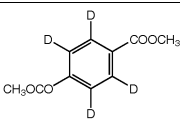
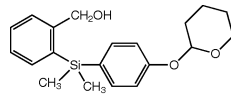
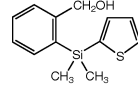
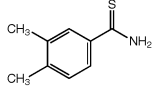
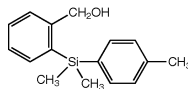
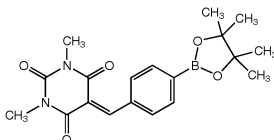
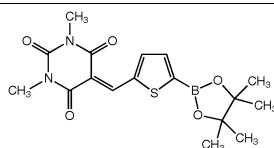
Stock #	Description		Size	\$
H32964	3,6-Dichloro-2-fluorophenol, JRD, 97% <chem>C6H3Cl2FO</chem> , F.W. 180.99, UN3261  R:21/22-34-51/53, S:26-36/37/39-45-61		250mg 1g	54.40 182.00
45607	Dichloro(mesitylene)ruthenium(II) dimer, Ru 34.6% [52462-31-4], <chem>C19H24Cl4Ru2</chem> , F.W. 584.34, m.p. 250° dec.			POR
45565	Dichloro(pentamethylcyclopentadienyl)iridium(III) dimer [12354-84-6], <chem>C20H30Cl4Ir2</chem> , F.W. 796.67, MDL MFCD00075435  R:36/37/38, S:26-37		500mg 2g	110.00 345.00
H30427	2,6-Dichloropyridine N-oxide, 98% [2587-00-0], <chem>C5H3Cl2NO</chem> , F.W. 163.99, m.p. 136-140°, UN2811  R:20/21/22-36/37/38, S:26-36/37		1g 5g	28.50 95.70
H31767	2,3-Dichloro-6-(trifluoromethyl)aniline, JRD, 97% <chem>C7H4Cl2F3N</chem> , F.W. 230.01, UN2811  R:20/21/22-33-36/38, S:26-36/37		250mg 1g	54.40 182.00
H32462	2,6-Dichloro-3-(trifluoromethyl)benzaldehyde, JRD, 97%  <chem>C8H3Cl2F3O</chem> , F.W. 243.01  R:36/37/38, S:26-37		1g 5g	75.60 305.00
H31956	2,6-Dichloro-3-(trifluoromethyl)benzoic acid, JRD, 97% <chem>C8H3Cl2F3O2</chem> , F.W. 259.01  R:36/37/38, S:26-37		1g 5g	62.70 250.00
H31792	2,6-Dichloro-3-(trifluoromethyl)benzyl bromide, JRD, 97% <chem>C8H4BrCl2F3</chem> , F.W. 307.92, m.p. 69-72°, UN3261  R:34, S:26-36/37/39-45		1g 5g	122.00 490.00
H31622	2,3-Dichloro-6-(trifluoromethyl)phenylacetic acid, JRD, 97% <chem>C9H5Cl2F3O2</chem> , F.W. 273.04  R:36/37/38, S:26-37		250mg 1g	67.70 226.00
H32151	2,6-Dichloro-3-(trifluoromethyl)phenylacetic acid, JRD, 97% <chem>C9H5Cl2F3O2</chem> , F.W. 273.04  R:36/37/38, S:26-37		250mg 1g	45.90 153.00
H32978	2,3-Dichloro-6-(trifluoromethyl)phenylacetonitrile, JRD, 97% <chem>C9H4Cl2F3N</chem> , F.W. 254.04, UN3276  R:20/21/22-36/38, S:26-36/37		250mg 1g	54.40 182.00

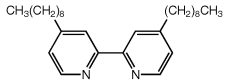
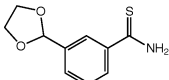
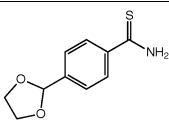
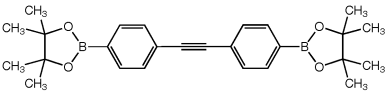
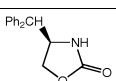
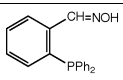
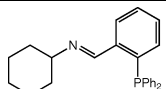
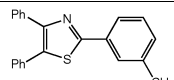
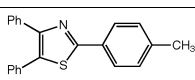
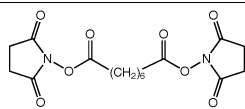
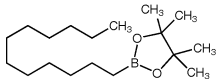
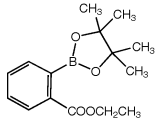
Stock #	Description		Size	\$
H51134	[2-(Dicyclohexylphosphino)ethyl]trimethylammonium chloride, 98% [181864-78-8], C ₁₇ H ₃₅ ClNP, F.W. 319.89, MDL MFCD05664248 R:36/37/38, S:26-37		1g 5g	99.80 394.00
H51694	6-(Diethoxymethyl)uracil, 98% [16953-48-3], C ₉ H ₁₄ N ₂ O ₄ , F.W. 214.22		250mg 1g	49.80 180.00
H51874	Diethyl (3-aminopropyl)phosphonate, 95%  [4402-24-8], C ₇ H ₁₈ NO ₃ P, F.W. 195.20, d. 1.053, UN2735, EINECS 224-535-6 R:34, S:26-36/37/39-45		250mg 1g	79.00 267.00
H32538	3,5-Diethyl-1,2-dihydro-1-phenyl-2-n-propylpyridine, tech. 75% [34562-31-7], C ₁₈ H ₂₅ N, F.W. 255.40, b.p. 125°/0.5mm, f.p. >100°(212°F), d. 0.97, n _D ²⁰ 1.5700, EINECS 252-091-3 R:36/38, S:23-26-37		10g 50g	102.00 338.00
H31544	Diethyl 4,4-difluoropimelate, 97% [22515-16-8], C ₁₁ H ₁₈ F ₂ O ₄ , F.W. 252.26, b.p. 108°/3mm, n _D ²⁰ 1.4150, MDL MFCD08146634		250mg 1g	99.20 277.00
H32891	Diethyl 4-fluoropimelate, 97% C ₁₁ H ₁₈ FO ₄ , F.W. 234.27, b.p. 86-87°/0.6mm, MDL MFCD08704574		250mg 1g	60.60 168.00
H51873	Diethyl [3-(N-phthalimido)propyl]phosphonate, 97% C ₁₅ H ₂₀ NO ₃ P, F.W. 325.30		1g 5g	45.00 191.00
H51145	1,3-Difluoroacetone, 98% [453-14-5], C ₃ H ₄ F ₂ O, F.W. 94.06, b.p. 101-102°, UN2929, MDL MFCD07368665 R:10-23/24/25, S:27-36/37-45		1g 5g	83.70 334.00
H32083	2,2-Difluorocyclopropanemethanol, 97% [509072-57-5], C ₃ H ₆ F ₂ O, F.W. 108.09, b.p. 129°, UN1987, MDL MFCD06797748 R:10-36/37/38, S:26-37		250mg 1g	171.00 476.00
H31817	2,2-Difluoro-N-(2-hydroxyethyl)propionamide, 98% CH ₃ CF ₂ CONH(CH ₂) ₂ OH, F.W. 153.13, b.p. 85°/0.2mm, MDL MFCD08687553 R:36/38, S:26-37		250mg 1g	121.00 336.00
H31816	3,5-Difluoro-2-methoxyaniline, JRD, 97% [41860-67-7], C ₇ H ₇ F ₂ NO, F.W. 159.14, UN2810 R:20/21/22-36/38, S:26-36/37		1g 5g	110.00 440.00
H31575	4,5-Difluoro-2-methoxyaniline, JRD, 97% [1017779-71-3], C ₇ H ₇ F ₂ NO, F.W. 159.14, UN2810 R:20/21/22-36/38, S:26-36/37		1g 5g	110.00 450.00
H51881	2,4-Difluoro-3-methoxybenzamide, 96% C ₈ H ₇ F ₂ NO ₂ , F.W. 187.14, m.p. 110-113° R:36/37/38, S:26-37		250mg 1g	107.00 345.00
H31980	3-(Difluoromethoxy)cinnamic acid, JRD, 97% C ₁₀ H ₈ F ₂ O ₃ , F.W. 214.17, m.p. 107-109° R:36/37/38, S:26-37		250mg 1g	54.40 182.00
H32283	3,5-Difluoro-2-methoxycinnamic acid, JRD, 97% C ₁₀ H ₈ F ₂ O ₃ , F.W. 214.17, m.p. 156-160° R:36/37/38, S:26-37		250mg 1g	54.40 182.00
H32170	4-(Difluoromethoxy)cinnamic acid, JRD, 97% C ₁₀ H ₈ F ₂ O ₃ , F.W. 214.17 R:36/37/38, S:26-37		250mg 1g	54.40 182.00

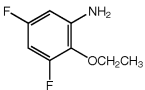
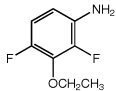
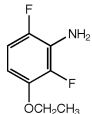
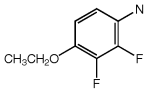
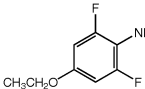
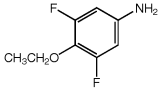
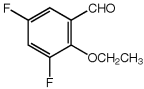
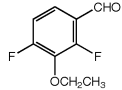
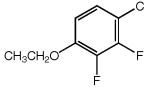
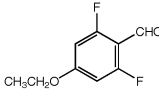
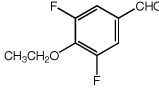
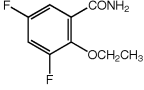
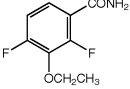
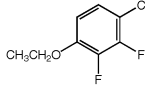
Stock #	Description		Size	\$
H32707	2-(Difluoromethoxy)-5-fluorobenzoic acid, JRD, 97% <chem>C8H5F3O3</chem> , F.W. 206.12 ✗ R:36/37/38, S:26-37		250mg 1g	67.70 230.00
H32600	4-Difluoromethoxy-2-fluorobenzoic acid, JRD, 97% <chem>C8H5F3O3</chem> , F.W. 206.12 ✗ R:36/37/38, S:26-37		250mg 1g	69.50 232.00
H31695	2-(Difluoromethoxy)-5-fluorobenzonitrile, JRD, 97% <chem>C8H4F3NO</chem> , F.W. 187.12, UN3276 ✗ R:20/21/22-36/38, S:26-36/37		250mg 1g	67.70 230.00
H31690	4-(Difluoromethoxy)-2-fluorobenzonitrile, JRD, 97% <chem>C8H4F3NO</chem> , F.W. 187.12, UN3439 ✗ R:20/21/22-36/37/38, S:26-36/37		250mg 1g	54.40 182.00
H32289	2-(Difluoromethoxy)-5-fluorobenzylamine, JRD, 97% △ <chem>C8H8F3NO</chem> , F.W. 191.15, UN2735 ✗ R:34, S:26-36/37/39-45		250mg 1g	81.70 273.00
H32849	4-(Difluoromethoxy)-3-methoxybenzaldehyde, JRD, 97% △ <chem>C9H8F2O3</chem> , F.W. 202.15 ✗ R:36/38, S:26-37		1g 5g	114.00 456.00
H31594	4-(Difluoromethoxy)-3-methoxybenzoic acid, JRD, 97% <chem>C9H8F2O4</chem> , F.W. 218.15 ✗ R:36/37/38, S:26-37		1g 5g	114.00 457.00
H31543	4-(Difluoromethoxy)-3-methoxycinnamic acid, JRD, 97% <chem>C11H10F2O4</chem> , F.W. 244.19 ✗ R:36/37/38, S:26-37		250mg 1g	54.40 182.00
H31511	3-[4-(Difluoromethoxy)-3-methoxyphenyl]propionic acid, JRD, 97% <chem>C11H12F2O4</chem> , F.W. 246.21 ✗ R:36/37/38, S:26-37		250mg 1g	67.70 226.00
H31728	2,6-Difluoro-3-methoxy-DL-phenylalanine, JRD, 97% <chem>C10H11F2NO3</chem> , F.W. 231.20		250mg 1g	67.70 226.00
H31574	3-(3,5-Difluoro-2-methoxyphenyl)propionic acid, JRD, 97% <chem>C10H10F2O3</chem> , F.W. 216.18 ✗ R:36/37/38, S:26-37		250mg 1g	67.70 226.00
H32688	3-(3,5-Difluoro-4-methoxyphenyl)propionic acid, JRD, 97% <chem>C10H10F2O3</chem> , F.W. 216.18 ✗ R:36/37/38, S:26-37		250mg 1g	67.70 225.00
H32399	3-(4,5-Difluoro-2-methoxyphenyl)propionic acid, JRD, 97% <chem>C10H10F2O3</chem> , F.W. 216.18 ✗ R:36/37/38, S:26-37		250mg 1g	67.70 230.00
H32938	3,5-Difluoro-2-methoxypyridine, 97% [1171918-06-1], <chem>C6H5F2NO</chem> , F.W. 145.11, m.p. 36-39° ✗ R:36/37/38, S:26-37		250mg 1g	59.90 166.00
H31989	3,4-Difluoro-5-methylbenzaldehyde, JRD, 97% △ <chem>C8H6F2O</chem> , F.W. 156.13 ✗ R:36/37/38, S:26-37		250mg 1g	54.40 182.00
H31851	3,5-Difluoro-2-methylbenzaldehyde, JRD, 97% △ <chem>C8H6F2O</chem> , F.W. 156.13, m.p. 32-34° ✗ R:36/37/38, S:26-37		250mg 1g	54.40 182.00

Stock #	Description		Size	\$
H32313	3,4-Difluoro-5-methylbenzoic acid, JRD, 97% C ₈ H ₆ F ₂ O ₂ , F.W. 172.13 R:36/37/38, S:26-37		250mg 1g	54.40 182.00
H31616	3,5-Difluoro-2-methylbenzoic acid, JRD, 97% C ₈ H ₆ F ₂ O ₂ , F.W. 172.13 R:36/37/38, S:26-37		250mg 1g	54.40 182.00
H32173	2,2-Difluoro-4-pentenoic acid, 97% [55039-89-9], C ₆ H ₈ F ₂ O ₂ , F.W. 136.10, b.p. 56-58°/3mm, UN3265, MDL MFCD09800642 R:34, S:23-26-36/37/39-45		250mg 1g 5g	50.40 140.00 448.00
H32976	2,3-Difluoro-4-n-pentylanisole, JRD, 97% C ₁₂ H ₁₈ F ₂ O, F.W. 214.25		250mg	198.00
H31740	2,3-Difluoro-4-n-pentylphenol, JRD, 97% C ₁₁ H ₁₄ F ₂ O, F.W. 200.23, UN3265 R:34, S:26-36/37/39-45		250mg	177.00
H32276	3,4-Difluoro-DL-phenylalanine, 97% [32133-36-1], C ₉ H ₉ F ₂ NO ₂ , F.W. 201.17, m.p. 237-240°		250mg 1g	43.90 146.00
H32408	2-(2,3-Difluorophenyl)ethanol, JRD, 97% C ₈ H ₈ F ₂ O, F.W. 158.15		250mg 1g	33.60 112.00
H32082	2-(2,5-Difluorophenyl)ethanol, JRD, 97% C ₈ H ₈ F ₂ O, F.W. 158.15		250mg 1g	33.60 112.00
H31710	2-(3,4-Difluorophenyl)ethanol, JRD, 97% C ₈ H ₈ F ₂ O, F.W. 158.15		250mg 1g	33.60 112.00
H32124	2,4-Difluoropyridine, 97% [34941-90-7], C ₅ H ₃ F ₂ N, F.W. 115.08, UN1993 R:10-36/37/38, S:26-37		250mg 1g 5g	71.20 197.00 660.00
H31130	2,5-Difluoropyridine, 97% [84476-99-3], C ₅ H ₃ F ₂ N, F.W. 115.08, UN1993 R:10-36/37/38, S:26-37		250mg 1g	46.20 129.00
H51068	(+)-Diisopropyl O,O'-bis(trimethylsilyl)-L-tartrate, 99% Δ [130678-42-1], C ₁₆ H ₃₄ O ₆ Si ₂ , F.W. 378.61, BRN 3620696, MDL MFCD00192055		250mg 1g 5g	41.90 121.00 492.00
H31707	2,4-Dimethoxybenzenesulfonamide, 96% [51770-71-9], C ₈ H ₁₁ NO ₃ S, F.W. 217.24, m.p. 166-168°, MDL MFCD06147001 R:36/37/38, S:26-37		1g 5g	106.00 354.00
H51086	2,2'-Dimethoxy-1,1'-binaphthyl, 97% [2960-93-2], C ₂₂ H ₁₈ O ₂ , F.W. 314.38, UN3077, MDL MFCD00091146 R:36/37/38-50/53, S:26-37-60-61		1g 5g	86.60 353.00
H51088	2,4-Dimethoxy-5-methylpyrimidine, 97% [5151-34-8], C ₇ H ₁₀ N ₂ O ₂ , F.W. 154.17, MDL MFCD03424272 R:36/37/38, S:26-37		1g	143.00

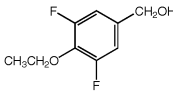
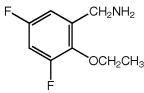
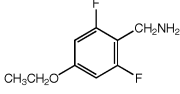
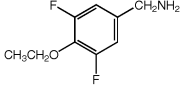
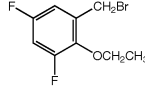
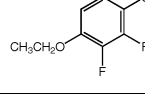
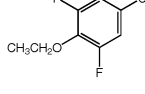
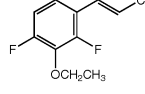
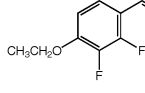
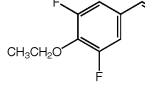
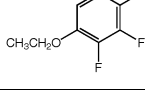
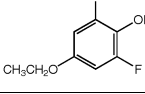
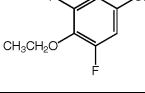
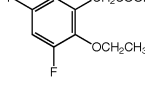
Stock #	Description		Size	\$
H32747	4,5-Dimethoxy-2-nitrophenylacetonitrile, 98% [17354-04-0], C ₁₀ H ₁₀ N ₂ O ₄ , F.W. 222.20, m.p. 110-112°, UN3439 ✗ R:20/21/22-36/37/38, S:26-36/37		250mg 1g	43.80 122.00
H51177	2-(2,5-Dimethoxyphenylamino)ethanol Δ [N-(2-Hydroxyethyl)-2,5-dimethoxyaniline] [28226-20-2], C ₁₀ H ₁₅ NO ₃ , F.W. 197.24, MDL MFCD03701227 ✗ R:36/37/38, S:26-37		250mg 1g	56.50 190.00
H51757	2-[(2,4-Dimethoxyphenyl)dimethylsilyl]benzyl alcohol, 95% C ₁₇ H ₂₂ O ₃ Si, F.W. 302.44, Note: Product of Advanced Molecular Technologies Pty Ltd. ✗ R:36/38, S:26-37		1g 5g	43.00 170.00
H51741	2-[(3,4-Dimethoxyphenyl)dimethylsilyl]benzyl alcohol, 95% C ₁₇ H ₂₂ O ₃ Si, F.W. 302.44, Note: Product of Advanced Molecular Technologies Pty Ltd. ✗ R:36/37/38, S:26-37		1g 5g	67.50 270.00
H51854	3,4-Dimethoxythiobenzamide, 97% [58952-14-0], C ₈ H ₁₁ NO ₂ S, F.W. 197.25, MDL MFCD04973329 ✗ R:22-36/37/38, S:26-36/37		250mg 1g	33.00 115.00
H51715	4-(Dimethylamino)pyridine, 99%, prilled [1122-58-3], C ₇ H ₁₀ N ₂ , F.W. 122.17, m.p. 110-113°, b.p. 162°/50mm, f.p. 124°(255°F), UN2928, EINECS 214-353-5, RTECS US9230000, BRN 110354, MDL MFCD00006418 ☠ R:24/25-34, S:26-36/37/39-45		5g 25g 100g	12.00 35.80 108.00
H30661	(S)-(-)-3-Dimethylaminopyrrolidine dihydrochloride, 97% ■ [(S)-(+)-N,N-Dimethyl-3-pyrrolidinamine dihydrochloride] [144043-20-9], C ₆ H ₁₄ N ₂ ·2HCl, F.W. 187.11, m.p. 176-181°, [α] _D ²⁰ -3.70° (c=1, water) ✗ R:36/37/38, S:26-37		250mg 1g	56.20 155.00
H51716	3,5-Dimethylbenzoyl chloride, 98% ▴ [6613-44-1], C ₉ H ₉ ClO, F.W. 168.62, b.p. 127°/20mm, d. 1.14, UN3265, RTECS DM6636900, MDL MFCD02313742 ☞ R:34-43, S:26-36/37/39-45		100g	108.00
H51139	N,N-Dimethyl-1,4-butanediamine dihydrochloride (CH ₃) ₂ NCH ₂ CH ₂ CH ₂ CH ₂ NH ₂ ·2HCl, F.W. 189.13, MDL MFCD11977712 ✗ R:22-36/37/38, S:26-36/37		1g 5g	48.60 182.00
H30194	3,3-Dimethyl-1-butanol, 97% [624-95-3], (CH ₃) ₂ CCCH ₂ CH ₂ OH, F.W. 102.18, m.p. -60°, b.p. 142-143°, f.p. 47°(117°F), d. 0.81, n _D ²⁰ 1.4140, UN1987, EINECS 210-872-6, t		10g 50g	81.60 280.00
H51739	2-(Dimethyl[4-(methylthio)phenyl]silyl)benzyl alcohol, 95% C ₁₈ H ₂₀ OSSi, F.W. 288.48, Note: Product of Advanced Molecular Technologies Pty Ltd. ✗ R:36/37/38, S:26-37		1g 5g	54.40 220.00
H51756	2-(Dimethyl[4-(4-morpholinyl)phenyl]silyl)benzyl alcohol, 95% C ₁₉ H ₂₅ NO ₂ Si, F.W. 327.49, Note: Product of Advanced Molecular Technologies Pty Ltd. ✗ R:36/37/38, S:26-37		1g 5g	69.40 275.00
H51696	2-[Dimethyl(1-naphthyl)silyl]benzyl alcohol, 97% C ₁₈ H ₂₀ OSi, F.W. 292.45, Note: Product of Advanced Molecular Technologies Pty Ltd. ✗ R:36/37/38, S:26-37		1g 5g	45.00 180.00
H51760	2-[Dimethyl(1-octenyl)silyl]benzyl alcohol, 95% [853955-61-0], C ₁₇ H ₂₈ O ₂ Si, F.W. 276.49, Note: Product of Advanced Molecular Technologies Pty Ltd. ✗ R:36/37/38, S:23-26-37		1g 5g	54.40 220.00
H30009	Dimethyl oxazole-4,5-dicarboxylate, 99% [Oxazole-4,5-dicarboxylic acid dimethyl ester] [72030-81-0], C ₇ H ₇ NO ₅ , F.W. 185.14, m.p. 74-76°, b.p. 150°/12mm		1g 10g	51.40 321.00

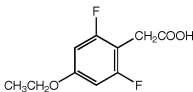
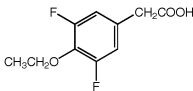
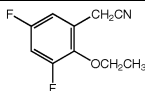
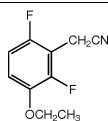
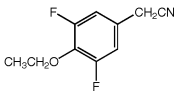
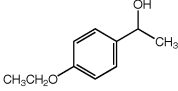
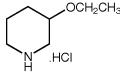
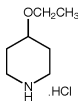
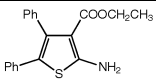
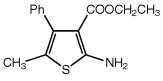
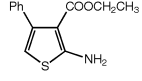
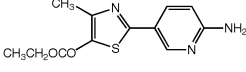
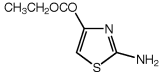
Stock #	Description		Size	\$
H51661	2-(Dimethylphenylsilyl)benzyl alcohol [853955-69-8], C ₁₅ H ₁₈ OSi, F.W. 242.39, MDL MFCD11870725, Note: Product of Advanced Molecular Technologies Pty Ltd.		1g	33.00
			5g	124.00
	☒ R:36/37/38, S:26-37			
H51783	2,5-Dimethyl-4-phenylthiazole, 97% C ₁₁ H ₁₁ NS, F.W. 189.28, m.p. 25-30°		1g	51.10
			5g	190.00
H51679	N,N-Dimethyl-2-(1-piperazinyl)nicotinamide hydrochloride, 96% C ₁₂ H ₁₈ N ₄ O·HCl, F.W. 270.76		250mg	64.50
			1g	232.00
	☒ R:36/37/38, S:26-37			
H51737	2-[Dimethyl(3-pyridyl)silyl]benzyl alcohol, 95% C ₁₄ H ₁₇ NOSi, F.W. 243.38, Note: Product of Advanced Molecular Technologies Pty Ltd.		1g	60.00
			5g	240.00
	☒ R:36/37/38, S:26-37			
H51759	2-[Dimethyl(β-styryl)silyl]benzyl alcohol, 95% [853955-63-2], C ₁₇ H ₂₀ OSi, F.W. 268.43, Note: Product of Advanced Molecular Technologies Pty Ltd.		1g	43.00
			5g	170.00
	☒ R:36/38, S:26-37			
H51104	Dimethyl terephthalate-d4(ring-d4), 95% [74079-01-9], C ₁₀ H ₆ D ₄ O ₄ , F.W. 198.21, MDL MFCD00182544		250mg	62.40
			1g	182.00
	☒ R:36/37/38, S:26-37			
H51758	2-(Dimethyl[4-(2-tetrahydropyranyloxy)phenyl]silyl)benzyl alcohol, 95% C ₂₀ H ₂₆ O ₅ Si, F.W. 342.50, Note: Product of Advanced Molecular Technologies Pty Ltd.		1g	61.90
			5g	248.00
	☒ R:36/38, S:26-37			
H51663	2-[Dimethyl(2-thienyl)silyl]benzyl alcohol [853955-72-3], C ₁₃ H ₁₆ OSSi, F.W. 248.42, MDL MFCD11870727, Note: Product of Advanced Molecular Technologies Pty Ltd.		1g	48.30
			5g	182.00
	☒ R:36/37/38, S:26-37			
H51793	3,4-Dimethylthiobenzamide, 97% [58952-03-7], C ₈ H ₁₁ NS, F.W. 165.25		250mg	74.00
			1g	254.00
	☒ R:22-36/37/38, S:26-36/37			
H31641	2,3-Dimethylthiophene, 97% [632-16-6], C ₆ H ₆ S, F.W. 112.19, b.p. 142°, f.p. 26° (79°F), UN1993		1g	92.40
			5g	308.00
	☒ R:10-20, S:16-36			
H51691	2-[Dimethyl(p-tolyl)silyl]benzyl alcohol, 97% C ₁₆ H ₂₀ OSi, F.W. 256.41, m.p. 55-57°, Note: Product of Advanced Molecular Technologies Pty Ltd.		1g	47.50
			5g	190.00
	☒ R:36/37/38, S:26-37			
H51124	4-(1,3-Dimethyl-2,4,6-trioxohexahydropyrimidin-5-yl-idenemethyl)benzeneboronic acid pinacol ester C ₁₉ H ₂₃ BN ₂ O ₅ , F.W. 370.21, MDL MFCD11113040		1g	40.00
			5g	153.00
H51898	2-(1,3-Dimethyl-2,4,6-trioxohexahydropyrimidin-5-yl-idenemethyl)thiophene-5-boronic acid pinacol ester, 95% C ₁₇ H ₂₁ BN ₂ O ₅ S, F.W. 376.23		1g	60.00
			5g	240.00

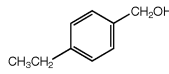
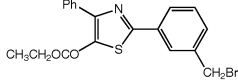
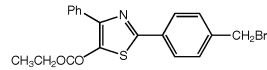
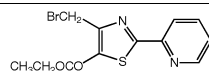
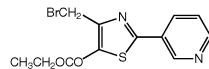
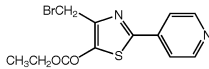
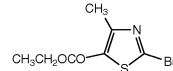
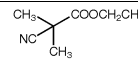
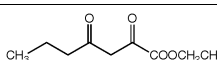
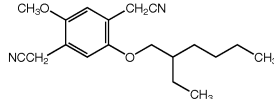
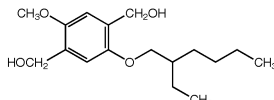
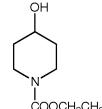
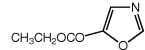
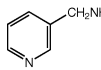
Stock #	Description	Size	\$
H32913	Dimethyl trisulfide, 98% [3658-80-8], CH ₃ SSSCH ₃ , F.W. 126.26, b.p. 165-175°, f.p. 52°(126°F), d. 1.203, n _D ²⁰ 1.600, UN1993, EINECS 222-910-9 ✗ R:10-36/37/38, S:23-26-37	5g 25g	83.70 280.00
H31476	4,4'-Di-n-nonyl-2,2'-bipyridine, 97% [4,4'-Dinonyl-2,2'-dipyridyl] [142646-58-0], C ₂₈ H ₄₄ N ₂ , F.W. 408.67, m.p. 61-63° ✗ R:36/37/38, S:26-37	 1g 5g	43.50 124.00
H51795	3-(1,3-Dioxolan-2-yl)thiobenzamide, 97% C ₁₀ H ₁₁ NO ₂ S, F.W. 209.26 ✗ R:22-36/37/38, S:26-36/37	 250mg 1g	74.00 254.00
H51856	4-(1,3-Dioxolan-2-yl)thiobenzamide, 97% [175202-43-4], C ₁₀ H ₁₁ NO ₂ S, F.W. 209.26, MDL MFCD00085174 ✗ R:22-36/37/38, S:26-36/37	 250mg 1g	74.00 254.00
H51897	Diphenylacetylene-4,4'-diboronic acid bis(pinacol) ester, 95% [849681-64-7], C ₂₆ H ₃₂ B ₂ O ₄ , F.W. 430.16	 1g 5g	78.80 310.00
H51074	(R)-(+)-4-(Diphenylmethyl)-2-oxazolidinone, 98% [173604-33-6], C ₁₆ H ₁₅ NO ₂ , F.W. 253.30, MDL MFCD01863565	 1g	190.00
H51903	2-(Diphenylphosphino)benzaldehyde oxime, 95% [153358-05-5], C ₁₉ H ₁₈ NOP, F.W. 305.32 ✗ R:36/37/38, S:26-37	 1g 5g	36.60 145.00
H51695	N-[2-(Diphenylphosphino)benzylidene]cyclohexylamine, 97% C ₂₈ H ₃₂ NP, F.W. 371.45 ✗ R:36/37/38, S:26-37	 1g 5g	120.00 490.00
H51805	4,5-Diphenyl-2-(m-tolyl)thiazole, 97% C ₂₂ H ₁₇ NS, F.W. 327.44 ✗ R:36/37/38, S:26-37	 250mg 1g	37.00 127.00
H51847	4,5-Diphenyl-2-(p-tolyl)thiazole, 97% C ₂₂ H ₁₇ NS, F.W. 327.44 ✗ R:36/37/38, S:26-37	 250mg 1g	37.00 127.00
H51771	Disuccinimidyl suberate, 97% [68528-80-3], C ₁₈ H ₂₀ N ₂ O ₈ , F.W. 368.34, m.p. 166-170°, MDL MFCD00049059	 500mg 1g	59.90 92.60
H31186	1-Dodecylboronic acid pinacol ester, 98% [2-n-Dodecyl-4,4,5,5-tetramethyl[1,3,2]dioxaborolane] [177035-82-4], C ₁₈ H ₃₇ BO ₂ , F.W. 296.31, b.p. 123°/0.2mm	 1g 5g 25g	54.10 180.00 596.00
H51172	1-Dodecylmagnesium bromide, 0.5M in MeTHF   [15890-72-9], CH ₃ (CH ₂) ₁₁ MgBr, F.W. 273.54, UN2924  R:11-14-19-34, S:8-16-26-36/37/39-43-45	100ml 500ml	30.70 115.00
H30347	Dowtherm® A [Diphenyl, Biphenyl Diphenyl ether eutectic] [8004-13-5], m.p. 12-14°, b.p. 257°, f.p. 113°, d. 1.056, UN3082, RTECS DV1500000, † ✗  R:36/37/38-50/53, S:26-37-57	1L 2.5L	84.00 168.00
H30941	2-(Ethoxycarbonyl)benzeneboronic acid pinacol ester, 97% [2-(Ethoxycarbonyl)phenylboronic acid pinacol ester, 2-(4,4,5,5-Tetramethyl-1,3,2-dioxaborolan-2-yl)benzoic acid ethyl ester] [269409-99-6], C ₁₅ H ₂₁ BO ₄ , F.W. 276.14, m.p. 57-62°	 1g 5g	90.10 348.00

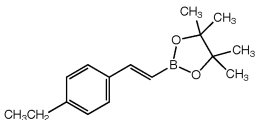
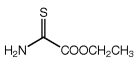
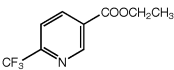
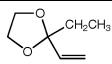
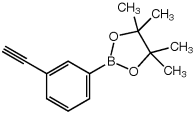
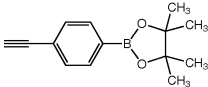
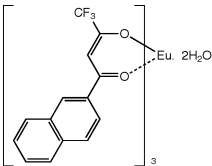
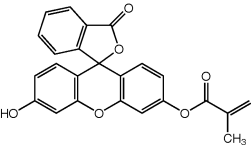
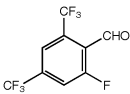
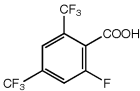
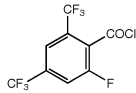
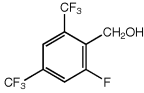
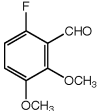
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H31965	3-Ethoxy-2,4-difluoroaniline, JRD, 97% C ₉ H ₇ F ₂ NO, F.W. 173.16, UN2810 ☒ R:20/21/22-36/38, S:26-28-36/37		1g 5g	110.00 440.00
H32732	3-Ethoxy-2,6-difluoroaniline, JRD, 97% C ₉ H ₇ F ₂ NO, F.W. 173.16, UN2810 ☒ R:20/21/22-36/38, S:26-28-36/37		1g 5g	122.00 490.00
H32627	4-Ethoxy-2,3-difluoroaniline, JRD, 97% C ₉ H ₇ F ₂ NO, F.W. 173.16, UN2810 ☒ R:20/21/22-36/38, S:26-28-36/37		1g 5g	95.70 383.00
H31722	4-Ethoxy-2,6-difluoroaniline, JRD, 97% C ₉ H ₇ F ₂ NO, F.W. 173.16, UN2810 ☒ R:20/21/22-36/38, S:26-28-36/37		250mg	170.00
H32031	4-Ethoxy-3,5-difluoroaniline, JRD, 97% C ₉ H ₇ F ₂ NO, F.W. 173.16, UN2811 ☒ R:20/21/22-36/38, S:26-36/37		1g 5g	110.00 439.00
H32294	2-Ethoxy-3,5-difluorobenzaldehyde, JRD, 97% △ C ₉ H ₆ F ₂ O ₂ , F.W. 186.16 ☒ R:36/38, S:26-37		1g 5g	78.10 312.00
H32631	3-Ethoxy-2,4-difluorobenzaldehyde, JRD, 97% △ C ₉ H ₆ F ₂ O ₂ , F.W. 186.16 ☒ R:36/38, S:26-37		1g 5g	45.30 182.00
H32450	4-Ethoxy-2,3-difluorobenzaldehyde, JRD, 97% △ C ₉ H ₆ F ₂ O ₂ , F.W. 186.16, m.p. 70-73° ☒ R:36/37/38, S:26-37		1g 5g	45.30 182.00
H31753	4-Ethoxy-2,6-difluorobenzaldehyde, JRD, 97% △ C ₉ H ₆ F ₂ O ₂ , F.W. 186.16, m.p. 70-72° ☒ R:36/37/38, S:26-37		1g 5g	74.40 298.00
H32671	4-Ethoxy-3,5-difluorobenzaldehyde, JRD, 97% △ C ₉ H ₆ F ₂ O ₂ , F.W. 186.16, m.p. 27-29° ☒ R:36/37/38, S:26-37		1g 5g	78.10 310.00
H32394	2-Ethoxy-3,5-difluorobenzamide, JRD, 97% C ₉ H ₆ F ₂ NO ₂ , F.W. 201.17 ☒ R:36/37/38, S:26-37		1g 5g	88.60 355.00
H32715	3-Ethoxy-2,4-difluorobenzamide, JRD, 97% C ₉ H ₆ F ₂ NO ₂ , F.W. 201.17 ☒ R:36/37/38, S:26-37		1g 5g	25.60 105.00
H31804	4-Ethoxy-2,3-difluorobenzamide, JRD, 97% C ₉ H ₆ F ₂ NO ₂ , F.W. 201.17 ☒ R:36/37/38, S:26-37		1g 5g	25.60 102.00

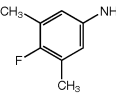
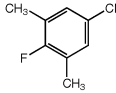
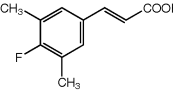
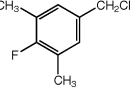
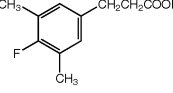
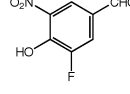
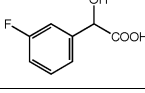
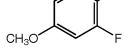
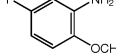
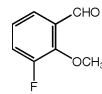
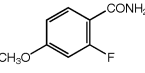
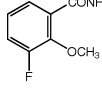
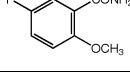
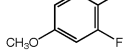
Stock #	Description		Size	\$
H32148	4-Ethoxy-2,6-difluorobenzamide, JRD, 97% C ₉ H ₈ F ₂ NO ₂ , F.W. 201.17 ✗ R:36/37/38, S:26-37		250mg	140.00
H32702	4-Ethoxy-3,5-difluorobenzamide, JRD, 97% C ₉ H ₈ F ₂ NO ₂ , F.W. 201.17 ✗ R:36/37/38, S:26-37		1g 5g	88.60 360.00
H32303	2-Ethoxy-3,5-difluorobenzoic acid, JRD, 97% C ₉ H ₈ F ₂ O ₃ , F.W. 202.15 ✗ R:36/37/38, S:26-37		1g 5g	78.10 312.00
H31812	3-Ethoxy-2,4-difluorobenzoic acid, JRD, 97% C ₉ H ₈ F ₂ O ₃ , F.W. 202.15 ✗ R:36/37/38, S:26-37		1g 5g	45.30 182.00
H31936	3-Ethoxy-2,6-difluorobenzoic acid, JRD, 97% C ₉ H ₈ F ₂ O ₃ , F.W. 202.15 ✗ R:36/37/38, S:26-37		1g 5g	75.60 303.00
H31849	4-Ethoxy-3,5-difluorobenzoic acid, JRD, 97% C ₉ H ₈ F ₂ O ₃ , F.W. 202.15 ✗ R:36/37/38, S:26-37		1g 5g	78.10 312.00
H32076	3-Ethoxy-2,4-difluorobenzonitrile, JRD, 97% C ₉ H ₇ F ₂ NO, F.W. 183.15, UN3276 ✗ R:20/21/22-36/38, S:26-36/37		1g 5g	110.00 439.00
H32114	3-Ethoxy-2,6-difluorobenzonitrile, JRD, 97% C ₉ H ₇ F ₂ NO, F.W. 183.15, UN3439 ✗ R:20/21/22-36/37/38, S:26-36/37		1g 5g	122.00 485.00
H31939	4-Ethoxy-2,3-difluorobenzonitrile, JRD, 97% C ₉ H ₇ F ₂ NO, F.W. 183.15, UN3439 ✗ R:20/21/22-36/37/38, S:26-36/37		1g 5g	110.00 439.00
H31715	4-Ethoxy-3,5-difluorobenzonitrile, JRD, 97% C ₉ H ₇ F ₂ NO, F.W. 183.15, UN3439 ✗ R:20/21/22-36/37/38, S:26-36/37		1g 5g	88.60 355.00
H31648	4-Ethoxy-2,3-difluorobenzoyl chloride, JRD, 97% C ₉ H ₇ ClF ₂ O ₂ , F.W. 220.60, UN3261 R:34, S:26-36/37/39-45		1g 5g	51.10 205.00
H32166	2-Ethoxy-3,5-difluorobenzyl alcohol, JRD, 97% C ₉ H ₁₀ F ₂ O ₂ , F.W. 188.17, n _D ²⁰ 1.4820 ✗ R:36/37/38, S:26-37		1g 5g	99.20 397.00
H32868	4-Ethoxy-2,3-difluorobenzyl alcohol, JRD, 97% C ₉ H ₁₀ F ₂ O ₂ , F.W. 188.17, m.p. 52-56° ✗ R:36/37/38, S:26-37		1g 5g	74.40 295.00
H32920	4-Ethoxy-2,6-difluorobenzyl alcohol, JRD, 97% C ₉ H ₁₀ F ₂ O ₂ , F.W. 188.17 ✗ R:36/37/38, S:26-37		250mg 1g	33.60 112.00

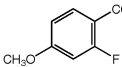
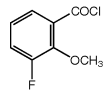
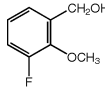
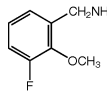
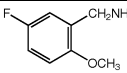
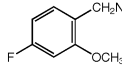
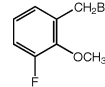
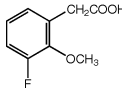
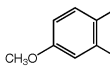
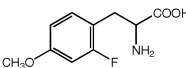
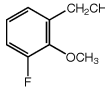
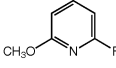
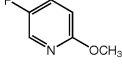
Stock #	Description		Size	\$
H32945	4-Ethoxy-3,5-difluorobenzyl alcohol, JRD, 97% C ₉ H ₁₀ F ₂ O ₂ , F.W. 188.17  R:36/37/38, S:26-37		1g 5g	110.00 440.00
H32972	2-Ethoxy-3,5-difluorobenzylamine, JRD, 97%  C ₉ H ₁₁ F ₂ NO, F.W. 187.19, UN2735  R:34, S:26-36/37/39-45		250mg 1g	67.70 230.00
H32007	4-Ethoxy-2,6-difluorobenzylamine, JRD, 97%  C ₉ H ₁₁ F ₂ NO, F.W. 187.19, UN2735  R:34, S:26-36/37/39-45		250mg 1g	86.80 289.00
H31875	4-Ethoxy-3,5-difluorobenzylamine, JRD, 97%  C ₉ H ₁₁ F ₂ NO, F.W. 187.19, UN2735  R:34, S:26-36/37/39-45		250mg 1g	67.70 225.00
H32126	2-Ethoxy-3,5-difluorobenzyl bromide, JRD, 97% C ₉ H ₉ BrF ₂ O, F.W. 251.07, UN3265  R:34, S:26-36/37/39-45		1g 5g	110.00 439.00
H32188	4-Ethoxy-2,3-difluorobenzyl bromide, JRD, 97% C ₉ H ₉ BrF ₂ O, F.W. 251.07, m.p. 58-61°, UN3261  R:34, S:26-36/37/39-45		1g 5g	95.70 383.00
H32801	4-Ethoxy-3,5-difluorobenzyl bromide, JRD, 97% C ₉ H ₉ BrF ₂ O, F.W. 251.07, UN3265  R:34, S:26-36/37/39-45		250mg 1g	54.40 182.00
H32160	3-Ethoxy-2,4-difluorocinnamic acid, JRD, 97% C ₁₁ H ₁₀ F ₂ O ₃ , F.W. 228.19  R:36/37/38, S:26-37		250mg 1g	47.60 159.00
H31772	4-Ethoxy-2,3-difluorocinnamic acid, JRD, 97% C ₁₁ H ₁₀ F ₂ O ₃ , F.W. 228.19  R:36/37/38, S:26-37		250mg 1g	40.80 136.00
H32893	4-Ethoxy-3,5-difluorocinnamic acid, JRD, 97% C ₁₁ H ₁₀ F ₂ O ₃ , F.W. 228.19  R:36/37/38, S:26-37		250mg 1g	47.60 159.00
H32029	4-Ethoxy-2,3-difluorophenol, JRD, 97% [126163-56-2], C ₉ H ₈ F ₂ O ₂ , F.W. 174.14  R:22-36/37/38, S:26-36/37		1g 5g	116.00 467.00
H32677	4-Ethoxy-2,6-difluorophenol, JRD, 97% C ₉ H ₈ F ₂ O ₂ , F.W. 174.14  R:22-36/37/38, S:26-36/37		250mg 1g	61.30 205.00
H31866	4-Ethoxy-3,5-difluorophenol, JRD, 97% C ₉ H ₈ F ₂ O ₂ , F.W. 174.14  R:22-36/37/38, S:23-26-36/37		250mg 1g	47.60 155.00
H31632	2-Ethoxy-3,5-difluorophenylacetic acid, JRD, 97% C ₁₀ H ₁₀ F ₂ O ₃ , F.W. 216.18  R:36/37/38, S:26-37		250mg 1g	81.70 275.00

Stock #	Description		Size	\$
H31592	4-Ethoxy-2,6-difluorophenylacetic acid, JRD, 97% C ₁₀ H ₁₀ F ₂ O ₃ , F.W. 216.18 ✗ R:36/37/38, S:26-37		250mg 1g	81.70 273.00
H31873	4-Ethoxy-3,5-difluorophenylacetic acid, JRD, 97% C ₁₀ H ₁₀ F ₂ O ₃ , F.W. 216.18 ✗ R:36/37/38, S:26-37		250mg 1g	81.70 273.00
H32578	2-Ethoxy-3,5-difluorophenylacetonitrile, JRD, 97% C ₁₀ H ₉ F ₂ NO, F.W. 197.18, UN3276 ✗ R:20/21/22-36/38, S:26-36/37		250mg 1g	67.70 225.00
H31931	3-Ethoxy-2,6-difluorophenylacetonitrile, JRD, 97% C ₁₀ H ₉ F ₂ NO, F.W. 197.18, m.p. 50-53°, UN3439 ✗ R:20/21/22-36/37/38, S:26-36/37		250mg 1g	54.40 182.00
H32781	4-Ethoxy-3,5-difluorophenylacetonitrile, JRD, 97% C ₁₀ H ₉ F ₂ NO, F.W. 197.18, UN3439 ✗ R:20/21/22-36/37/38, S:26-36/37		250mg 1g	67.70 230.00
H51865	1-Ethoxymethyl-2-iodoimidazole ▲ [146697-87-2], C ₆ H ₉ IN ₂ O, F.W. 252.05 ✗ R:36/38, S:26-37		250mg 1g	51.00 150.00
H30052	1-(4-Ethoxyphenyl)ethanol, 98% [52067-36-4], C ₁₀ H ₁₄ O ₂ , F.W. 166.22, m.p. 46-47°		1g 5g 25g	26.90 89.90 300.00
H51152	3-Ethoxypiperidine hydrochloride C ₇ H ₁₃ NO·HCl, F.W. 165.66, MDL MFCD09475416 ✗ R:36/37/38, S:26-37		250mg 1g	57.60 208.00
H51028	4-Ethoxypiperidine hydrochloride C ₇ H ₁₃ NO·HCl, F.W. 165.66, MDL MFCD06804554 ✗ R:36/37/38, S:26-37		250mg 1g	59.40 214.00
H51846	Ethyl 2-amino-4,5-diphenylthiophene-3-carboxylate, 97% C ₁₉ H ₁₇ NO ₂ S, F.W. 323.41 ✗ R:36/37/38, S:26-37		1g 5g	88.00 328.00
H51799	Ethyl 2-amino-5-methyl-4-phenylthiophene-3-carboxylate, 97% [4815-37-6], C ₁₄ H ₁₅ NO ₂ S, F.W. 261.34, MDL MFCD01763475 ✗ R:36/38, S:26-37		1g 5g	79.00 294.00
H51732	Ethyl 2-amino-4-phenylthiophene-3-carboxylate, 97% [4815-36-5], C ₁₃ H ₁₃ NO ₂ S, F.W. 247.42, m.p. 76-78°, RTECS XM8350000 ✗ R:36/37/38, S:26-37		1g 5g	75.80 276.00
H51842	Ethyl 2-(2-amino-5-pyridyl)-4-methylthiazole-5-carboxylate, 97% C ₁₂ H ₁₃ N ₃ O ₂ S, F.W. 263.31 ✗ R:36/37/38, S:26-37		250mg 1g	128.00 384.00
H30634	Ethyl 2-aminothiazole-4-carboxylate, 98% [5398-36-7], C ₆ H ₈ N ₂ O ₂ S, F.W. 172.21, m.p. 175-178° ✗ R:36/37/38, S:26-37		1g 10g	28.90 240.00
H51682	Ethyl 5-aminovalerate hydrochloride, 96% [29840-57-1], H ₂ N(CH ₂) ₄ CO ₂ CH ₂ CH ₃ ·HCl, F.W. 181.66 ✗ R:36/37/38, S:26-37		250mg 1g	53.20 192.00

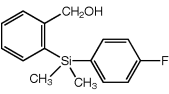
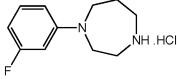
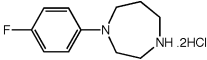
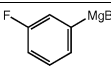
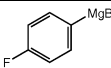
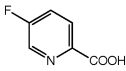
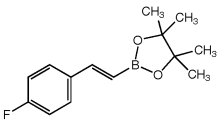
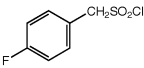
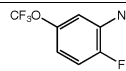
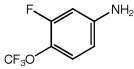
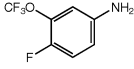
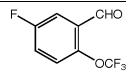
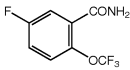
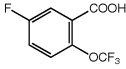
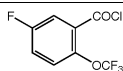
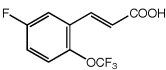
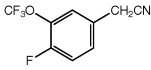
Stock #	Description		Size	\$
H30459	4-Ethylbenzyl alcohol, 99% [768-59-2], C ₉ H ₁₀ O, F.W. 136.19, b.p. 115-117°/9mm, d. 1.028, n _D ²⁰ 1.5275, EINECS 292-198-8, MDL MFCD00004666		1g 5g	46.60 140.00
H51820	Ethyl 2-[3-(bromomethyl)phenyl]-4-phenyl-thiazole-5-carboxylate, 97% C ₁₉ H ₁₆ BrNO ₂ S, F.W. 402.31, UN3261  R:34, S:26-36/37/39-45		250mg 1g	37.00 127.00
H51813	Ethyl 2-[4-(bromomethyl)phenyl]-4-phenyl-thiazole-5-carboxylate, 97% C ₁₉ H ₁₆ BrNO ₂ S, F.W. 402.31, UN3261  R:34, S:26-36/37/39-45		250mg 1g	37.00 127.00
H51727	Ethyl 4-bromomethyl-2-(2-pyridyl)thiazole-5-carboxylate, 97% [1138444-37-7], C ₁₂ H ₁₁ BrN ₂ O ₂ S, F.W. 327.20, m.p. 100-104°, UN3261  R:34, S:26-36/37/39-45		1g 5g	71.30 260.00
H51811	Ethyl 4-bromomethyl-2-(3-pyridyl)thiazole-5-carboxylate, 95% C ₁₂ H ₁₁ BrN ₂ O ₂ S, F.W. 327.20, UN3261  R:34, S:26-36/37/39-45		1g 5g	71.00 266.00
H51817	Ethyl 4-bromomethyl-2-(4-pyridyl)thiazole-5-carboxylate, 95% C ₁₂ H ₁₁ BrN ₂ O ₂ S, F.W. 327.20, UN3261  R:34, S:26-36/37/39-45		1g 5g	71.00 266.00
H51833	Ethyl 2-bromo-4-methylthiazole-5-carboxylate, 97% [22900-83-0], C ₇ H ₈ BrNO ₂ S, F.W. 250.12, MDL MFCD03791227  R:36/37/38, S:26-37		1g 5g	50.00 185.00
H31432	Ethyl 2-cyano-2-methylpropionate, 97% [2-Cyano-2-methylpropionic acid ethyl ester, Ethyl cyanodimethylacetate] [1572-98-1], C ₇ H ₁₁ NO ₂ , F.W. 141.17, b.p. 185°, d. 0.97  R:22, S:36		10g 50g	64.20 214.00
H31027	Ethyl 2,4-dioxoheptanoate, 95% [2,4-Dioxoheptanoic acid ethyl ester] [36983-31-0], C ₉ H ₁₄ O ₄ , F.W. 186.21, b.p. 228-232°		1g 5g 25g	23.10 91.90 369.00
H31852	Ethyl 2,2-difluoropropionate, 97% [28781-85-3], CH ₃ CF ₂ CO ₂ CH ₂ CH ₃ , F.W. 138.11, b.p. 24°/20mm, UN3272, MDL MFCD06657970  R:10-36/37/38, S:23-26-37		1g 5g	69.00 231.00
H51040	2-(2-Ethylhexyloxy)-5-methoxy-1,4-benzenediacetonitrile, 98% [213749-91-8], C ₁₉ H ₂₆ N ₂ O ₂ , F.W. 314.42, UN3439, MDL MFCD03701544  R:20/21/22-36/37/38, S:26-36/37		1g 5g	148.00 604.00
H51042	2-(2-Ethylhexyloxy)-5-methoxy-1,4-benzenedimethanol, 98%  [245731-58-2], C ₁₇ H ₂₈ O ₄ , F.W. 296.40, MDL MFCD03427231		1g 5g	113.00 459.00
H31388	Ethyl 4-hydroxypiperidine-1-carboxylate, 98% [4-Hydroxypiperidine-1-carboxylic acid ethyl ester, 1-Ethoxycarbonyl-1-piperidinol] [65214-82-6], C ₈ H ₁₅ NO ₃ , F.W. 173.21, n _D ²⁰ 1.4802, EINECS 265-636-5, MDL MFCD00086880		5g 25g 100g	52.40 174.00 483.00
H31372	Ethyl isethionate, 95% [Isethionic acid ethyl ester, Ethyl 2-hydroxyethanesulfonate] [58337-44-3], HO(CH ₂) ₂ SO ₃ CH ₂ CH ₃ , F.W. 154.19  R:36/37/38, S:26-37		1g 10g	49.80 310.00
H32924	Ethyl oxazole-5-carboxylate, 98% [118994-89-1], C ₆ H ₇ NO ₃ , F.W. 141.13, b.p. 60°/2.25mm, n _D ²⁰ 1.4680, BRN 83084416		5g 25g	67.20 224.00
H51120	N-Ethyl-3-pyridinemethylamine hydrochloride C ₈ H ₁₂ N ₂ ·HCl, F.W. 172.66, MDL MFCD07106861  R:36/37/38		250mg 1g	62.60 202.00

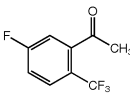
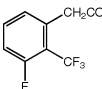
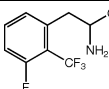
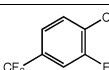
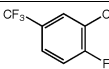
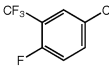
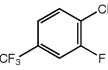
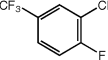
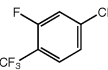
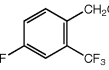
Stock #	Description		Size	\$
H30744	4-Ethyl-trans-β-styrylboronic acid pinacol ester, 97% [2-[trans-2-(4-Ethylphenyl)vinyl]-4,4,5,5-tetramethyl- [1,3,2]dioxaborolane] [870717-91-2], C ₁₆ H ₂₃ BO ₂ , F.W. 258.17, m.p. 46-48°, f.p. >110° (230°F)		250mg	39.50
			1g	110.00
			5g	367.00
H31447	Ethyl thiooxamate, 95% [Ethyl aminothioxoacetate, Thiooxamic acid ethyl ester] [16982-21-1], C ₄ H ₇ NO ₂ S, F.W. 133.17, m.p. 61-65° ✗ R:36/37/38, S:26-37		1g	31.00
			5g	102.00
			25g	337.00
H31896	Ethyl 6-(trifluoromethyl)nicotinate, 97% C ₉ H ₈ F ₃ NO ₂ , F.W. 219.16, MDL MFCD04972865 ✗ R:36/38, S:26-37		250mg	59.50
			1g	165.00
H31971	2-Ethyl-2-vinyl-1,3-dioxolane, 98% [22515-82-8], C ₇ H ₁₂ O ₂ , F.W. 128.17, b.p. 70°/77mm, UN3271 ✗ R:10-36, S:16-26		10g	61.40
			50g	205.00
H51902	3-Ethynylbenzeneboronic acid pinacol ester, 95% C ₁₄ H ₁₇ BO ₂ , F.W. 228.09 ✗ R:36/37/38, S:26-37		1g	88.10
			5g	350.00
H51745	4-Ethynylbenzeneboronic acid pinacol ester, 95% [1034287-04-1], C ₁₄ H ₁₇ BO ₂ , F.W. 228.09		1g	62.00
			5g	260.00
H32211	Europium(III) 1-(2-naphthyl)-4,4,4-trifluoro-1,3-butanedionate dihydrate C ₄₂ H ₂₇ EuF ₉ O ₉ ·2H ₂ O, F.W. 986.65 (950.62anhy), m.p. 126-132° ✗ R:36/37/38, S:26-37		250mg	51.50
			1g	143.00
H51046	Fluorescein O-methacrylate, 97% △ [480439-15-4], C ₂₄ H ₁₆ O ₆ , F.W. 400.38, MDL MFCD04974071 ✗ R:36/37/38, S:26-37		1g	105.00
			5g	427.00
H32020	2-Fluoro-4,6-bis(trifluoromethyl)benzaldehyde, JRD, 97% △ C ₉ H ₃ F ₇ O, F.W. 260.11 ✗ R:36/38, S:26-37		1g	78.10
			5g	312.00
H32718	2-Fluoro-4,6-bis(trifluoromethyl)benzoic acid, JRD, 97% C ₉ H ₃ F ₇ O ₂ , F.W. 276.11 ✗ R:36/37/38, S:26-37		1g	78.10
			5g	315.00
H32292	2-Fluoro-4,6-bis(trifluoromethyl)benzoyl chloride, JRD, 97% ▢ C ₉ H ₂ ClF ₇ O, F.W. 294.55, UN3265 ✗ R:34, S:26-36/37/39-45		1g	103.00
			5g	410.00
H31928	2-Fluoro-4,6-bis(trifluoromethyl)benzyl alcohol, JRD, 97% C ₉ H ₅ F ₇ O, F.W. 262.12, m.p. 32-34° ✗ R:36/38, S:26-37		250mg	40.80
			1g	136.00
H51877	6-Fluoro-2,3-dimethoxybenzaldehyde, 97% △ C ₉ H ₇ FO ₃ , F.W. 184.16, m.p. 45-47° ✗ R:36/37/38, S:26-37		250mg	95.00
			1g	306.00

Stock #	Description		Size	\$
H31663	4-Fluoro-3,5-dimethylaniline, JRD, 97% [4-Fluoro-3,5-xylidine] C ₈ H ₁₀ FN, F.W. 139.17, UN2811 ☒ R:20/21/22-33-36/38, S:26-28-36/37		250mg 1g	48.60 162.00
H31712	4-Fluoro-3,5-dimethylbenzonitrile, JRD, 97% C ₈ H ₈ FN, F.W. 149.06, UN3276 ☒ R:20/21/22-36/38, S:26-36/37		250mg 1g	47.60 159.00
H31984	4-Fluoro-3,5-dimethylcinnamic acid, JRD, 97% C ₁₁ H ₁₁ FO ₂ , F.W. 194.20 ☒ R:36/37/38, S:26-37		250mg 1g	47.60 159.00
H31797	4-Fluoro-3,5-dimethylphenylacetonitrile, JRD, 97% C ₁₀ H ₁₀ FN, F.W. 163.19, UN3276 ☒ R:20/21/22-36/38, S:26-36/37		250mg 1g	47.60 182.00
H31576	3-(4-Fluoro-3,5-dimethylphenyl)propionic acid, JRD, 97% C ₁₁ H ₁₃ FO ₂ , F.W. 196.22 ☒ R:36/37/38, S:26-37		250mg 1g	67.70 226.00
H51878	3-Fluoro-4-hydroxy-5-nitrobenzaldehyde, 95% △ C ₇ H ₄ FNO ₄ , F.W. 185.11 ☒ R:36/37/38, S:26-37		250mg 1g	47.00 153.00
H31270	3-Fluoromandelic acid, 97% [3-Fluoro-α-hydroxyphenylacetic acid] [395-05-1], C ₈ H ₇ FO ₃ , F.W. 170.14, m.p. 93-98° ☒ R:36/37/38, S:26-37		1g 5g	38.20 127.00
H32885	2-Fluoro-4-methoxyaniline, JRD, 97% [458-52-6], C ₇ H ₈ FNO, F.W. 141.14, m.p. 48-51°, UN2811 ☒ R:24/25-33-36/37/38, S:26-28-36/37-45		250mg 1g	41.80 140.00
H32214	3-Fluoro-2-methoxyaniline, JRD, 97% [437-83-2], C ₇ H ₈ FNO, F.W. 141.14, UN2810 ☒ R:24/25-33-36/38, S:26-28-36/37-45		250mg 1g	95.20 317.00
H32525	5-Fluoro-2-methoxyaniline, JRD, 97% [1978-39-8], C ₇ H ₈ FNO, F.W. 141.14, UN2810 ☒ R:24/25-33-36/38, S:26-28-36/37/39-45		250mg 1g	40.80 136.00
H32971	3-Fluoro-2-methoxybenzaldehyde, JRD, 97% △ [74266-68-5], C ₈ H ₇ FO ₂ , F.W. 154.14 ☒ R:36/37/38, S:26-37		1g 5g	25.60 105.00
H31636	2-Fluoro-4-methoxybenzamide, JRD, 97% C ₈ H ₈ FNO ₂ , F.W. 169.15 ☒ R:36/37/38, S:26-37		1g 5g	88.60 355.00
H31949	3-Fluoro-2-methoxybenzamide, JRD, 97% C ₈ H ₈ FNO ₂ , F.W. 169.15 ☒ R:36/37/38, S:26-37		250mg 1g	95.20 317.00
H31588	5-Fluoro-2-methoxybenzamide, JRD, 97% C ₈ H ₈ FNO ₂ , F.W. 169.15 ☒ R:36/37/38, S:26-37		1g 5g	88.60 355.00
H32531	2-Fluoro-4-methoxybenzonitrile, JRD, 97% [94610-82-9], C ₈ H ₇ FNO, F.W. 151.14, UN3439 ☒ R:20/21/22-36/37/38, S:26-36/37		1g 5g	103.00 410.00

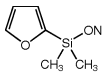
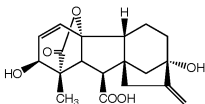
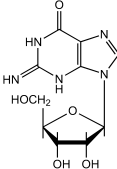
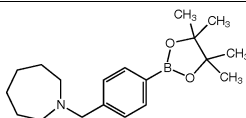
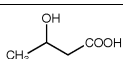
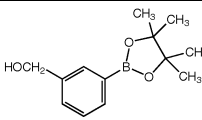
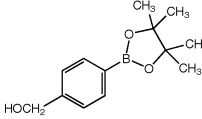
Stock #	Description		Size	\$
H32284	3-Fluoro-2-methoxybenzonitrile, JRD, 97% C ₈ H ₆ FNO, F.W. 151.14, UN3276  R:20/21/22-36/38, S:26-36/37		1g 5g	56.40 226.00
H31644	2-Fluoro-4-methoxybenzoyl chloride, JRD, 97%  C ₈ H ₆ ClFO ₂ , F.W. 188.58, UN3261  R:34, S:26-36/37/39-45		1g 5g	107.00 422.00
H32951	3-Fluoro-2-methoxybenzoyl chloride, JRD, 97%  C ₈ H ₆ ClFO ₂ , F.W. 188.58, UN3265  R:34, S:26-36/37/39-45		250mg 1g	81.70 273.00
H32416	3-Fluoro-2-methoxybenzyl alcohol, JRD, 97% C ₈ H ₈ FO ₂ , F.W. 156.15  R:36/37/38, S:23-26-37		250mg 1g	81.70 275.00
H32293	3-Fluoro-2-methoxybenzylamine, JRD, 97%  C ₈ H ₁₀ FNO, F.W. 155.17, UN2735  R:34, S:26-36/37/39-45		250mg 1g	95.20 317.00
H32897	5-Fluoro-2-methoxybenzylamine, JRD, 97%  C ₈ H ₁₀ FNO, F.W. 155.17, UN2735  R:34, S:26-36/37/39-45		250mg 1g	67.70 226.00
H51153	4-Fluoro-2-methoxybenzylamine hydrochloride, 97% C ₈ H ₁₀ FNO·HCl, F.W. 191.63, MDL MFCD04116342  R:36/37/38, S:26-37		1g 5g	78.40 240.00
H32562	3-Fluoro-2-methoxybenzyl bromide, JRD, 97% C ₈ H ₈ BrFO, F.W. 219.05, UN3265  R:34, S:26-36/37/39-45		250mg 1g	95.20 317.00
H32701	3-Fluoro-2-methoxycinnamic acid, JRD, 97% C ₁₀ H ₈ FO ₃ , F.W. 196.18  R:36/37/38, S:26-37		250mg 1g	54.40 185.00
H31678	3-Fluoro-2-methoxyphenylacetic acid, JRD, 97% [1017778-30-1], C ₉ H ₈ FO ₃ , F.W. 184.16  R:36/37/38, S:26-37		250mg	128.00
H30752	2-Fluoro-4-methoxyphenylacetonitrile, 97% [4-Methoxy-2-fluorobenzyl cyanide] [749934-29-0], C ₈ H ₆ FNO, F.W. 165.17, n _D ²⁰ 1.5131, UN3276  R:20/21/22-36/37/38, S:26-36/37		250mg 1g	69.10 230.00
H31729	2-Fluoro-4-methoxy-DL-phenylalanine, JRD, 97% C ₁₀ H ₁₂ FNO ₃ , F.W. 213.21		250mg 1g	67.70 226.00
H31843	3-(3-Fluoro-2-methoxyphenyl)propionic acid, JRD, 97% C ₁₀ H ₁₁ FO ₃ , F.W. 198.19  R:36/37/38, S:26-37		250mg 1g	67.70 226.00
H32994	2-Fluoro-6-methoxypyridine, 97% [116241-61-3], C ₆ H ₆ FNO, F.W. 127.12, UN1993  R:10-36/37/38, S:23-26-37		250mg 1g	103.00 287.00
H31901	3-Fluoro-2-methoxypyridine, 97% [884494-69-3], C ₆ H ₆ FNO, F.W. 127.12, UN1993  R:10-36/37/38, S:23-26-37		250mg 1g	114.00 316.00
H30991	5-Fluoro-2-methoxypyridine, 97% [51173-04-7], C ₆ H ₆ FNO, F.W. 127.12, UN1993  R:10-36/37/38, S:23-26-37		1g 5g 25g	64.60 215.00 720.00

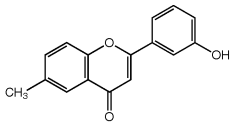
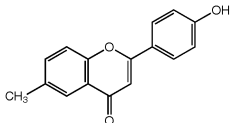
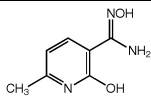
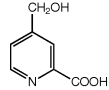
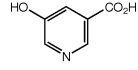
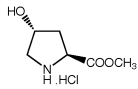
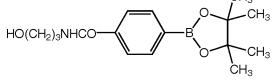
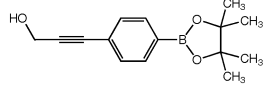
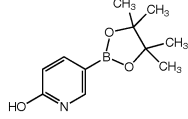
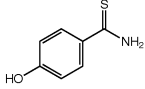
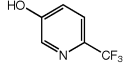
Stock #	Description		Size	\$
H32221	3'-Fluoro-2'-methylacetophenone, JRD, 97% C ₉ H ₈ FO, F.W. 152.17 R:36/38, S:26-37		1g 5g	103.00 411.00
H32499	4'-Fluoro-3'-methylacetophenone, JRD, 97% [369-32-4], C ₉ H ₈ FO, F.W. 152.17 R:36/38, S:26-37		1g 5g	34.00 136.00
H31292	2-Fluoro-4-methylbenzoyl chloride, 97% [59189-98-9], C ₈ H ₇ ClFO, F.W. 172.59, m.p. 43-47°, UN3261 R:34, S:26-36/37/39-45		1g 5g	88.90 355.00
H31864	3-Fluoro-4-methylphenol, JRD, 97% [452-78-8], C ₇ H ₇ FO, F.W. 126.13, UN3265 R:21/22-34, S:26-36/37/39-45		1g 5g	99.20 395.00
H51688	2-Fluoro-N-methyl-6-phenoxybenzylamine hydrochloride, 98% C ₁₄ H ₁₄ FNO·HCl, F.W. 267.73 R:36/37/38, S:26-37		250mg 1g	59.40 214.00
H30525	4-Fluoro-3-methylphenylacetic acid, 97% [1000520-92-2], C ₉ H ₈ FO ₂ , F.W. 168.17, m.p. 105-107° R:36/37/38, S:26-37		250mg 1g	37.30 125.00
H32846	3-Fluoro-2-methylphenylacetone nitrile, 97% [500912-15-2], C ₉ H ₈ FN, F.W. 149.17, UN3439 R:20/21/22-36/37/38, S:26-36/37		250mg 1g	40.80 137.00
H32194	3-Fluoro-4-methyl-DL-phenylalanine, JRD, 97% C ₁₀ H ₁₂ FNO ₂ , F.W. 197.21		250mg 1g	67.70 226.00
H32370	3-(2-Fluoro-4-methylphenyl)propionic acid, JRD, 97% C ₁₀ H ₁₁ FO ₂ , F.W. 182.19 R:36/37/38, S:26-37		250mg 1g	67.70 230.00
H32105	3-(3-Fluoro-4-methylphenyl)propionic acid, JRD, 97% C ₁₀ H ₁₁ FO ₂ , F.W. 182.19 R:36/37/38, S:26-37		250mg 1g	54.40 182.00
H31624	3-(4-Fluoro-2-methylphenyl)propionic acid, JRD, 97% C ₁₀ H ₁₁ FO ₂ , F.W. 182.19 R:36/37/38, S:26-37		250mg 1g	54.40 182.00
H32474	3-(4-Fluoro-3-methylphenyl)propionic acid, JRD, 97% C ₁₀ H ₁₁ FO ₂ , F.W. 182.19 R:36/37/38, S:26-37		250mg 1g	54.40 182.00
H32947	3-(5-Fluoro-2-methylphenyl)propionic acid, JRD, 97% C ₁₀ H ₁₁ FO ₂ , F.W. 182.19 R:36/37/38, S:26-37		250mg 1g	67.70 225.00
H31979	3-Fluoro-4-nitrobenzenesulfonyl chloride, 98% [86156-93-6], C ₈ H ₆ ClFNO ₃ S, F.W. 239.61, m.p. 44-48°, UN3261, MDL MFCD03412240 R:34, S:26-36/37/39-45		250mg 1g	60.60 168.00
H32176	5-Fluorooxindole, 97% [56341-41-4], C ₈ H ₆ FNO, F.W. 151.14, m.p. 142-146°, MDL MFCD02179598 R:36/37/38, S:26-37		1g	104.00
H51681	2-(3-Fluorophenoxy)ethylamine hydrochloride, 98% C ₉ H ₁₀ FNO·HCl, F.W. 191.63 R:36/37/38, S:26-37		250mg 1g	61.60 220.00
H31410	2-(4-Fluorophenoxy)ethylamine hydrochloride, 97% [263409-81-0], C ₉ H ₁₀ FNO·HCl, F.W. 191.63, m.p. 213-216° R:36/37/38, S:26-37		250mg 1g	81.60 293.00

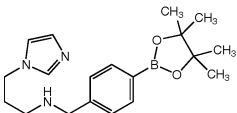
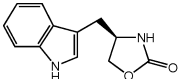
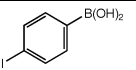
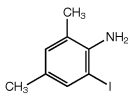
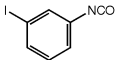
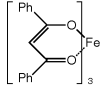
Stock #	Description	Size	\$
H51735	2-[(4-Fluorophenyl)dimethylsilyl]benzyl alcohol, 95% [853955-70-1], C ₁₅ H ₁₇ FOSi, F.W. 260.38, Note: Product of Advanced Molecular Technologies Pty Ltd. 	1g 5g	40.50 160.00
H51686	1-(3-Fluorophenyl)homopiperazine monohydrochloride, 98% [934991-99-8], C ₁₁ H ₁₅ FN ₂ ·HCl, F.W. 230.71, m.p. 187-191°, Note: Sold in collaboration with Tosoh Organic Chemical Co., Ltd. 	500mg	390.00
H51701	1-(4-Fluorophenyl)homopiperazine dihydrochloride, 98% [263409-96-7], C ₁₁ H ₁₅ FN ₂ ·2HCl, F.W. 230.71, m.p. 219-223°, Note: Sold in collaboration with Tosoh Organic Chemical Co., Ltd. 	500mg	390.00
H51168	3-Fluorophenylmagnesium bromide, 1M in MeTHF   [17318-03-5], C ₆ H ₄ BrFMg, F.W. 199.31, UN2924 	100ml 500ml	28.80 109.00
H51169	4-Fluorophenylmagnesium bromide, 1M in MeTHF   [352-13-6], C ₆ H ₄ BrFMg, F.W. 199.31, UN2924 	100ml 500ml	31.70 119.00
H32086	5-Fluoropyridine-2-carboxylic acid, 97% [107504-08-5], C ₆ H ₄ FNO ₂ , F.W. 141.10, m.p. 164-166° 	250mg 1g	82.80 230.00
H30578	4-Fluoro-trans-β-styrylboronic acid pinacol ester, 97% [2-[trans-2-(4-Fluorophenyl)vinyl]-4,4,5,5-tetramethyl-1,3,2]dioxaborolane] [504433-86-7], C ₁₄ H ₁₈ BFO ₂ , F.W. 248.11 	250mg 1g 5g	52.20 145.00 485.00
H31605	4-Fluoro-α-toluenesulfonyl chloride, 97%  [103360-04-9], C ₇ H ₆ ClFO ₂ S, F.W. 208.64, m.p. 58-61°, UN3261, MDL MFCD01631930 	250mg 1g	59.90 166.00
H32103	2-Fluoro-5-(trifluoromethoxy)aniline, JRD, 97% [116369-23-4], C ₇ H ₅ F ₄ NO, F.W. 195.11, UN2810 	1g 5g	117.00 467.00
H32561	3-Fluoro-4-(trifluoromethoxy)aniline, JRD, 97% C ₇ H ₅ F ₄ NO, F.W. 195.11, UN2810 	250mg 1g	81.70 273.00
H32810	4-Fluoro-3-(trifluoromethoxy)aniline, JRD, 97% C ₇ H ₅ F ₄ NO, F.W. 195.11, UN2810 	250mg 1g	110.00 365.00
H32956	5-Fluoro-2-(trifluoromethoxy)benzaldehyde, JRD, 97%  C ₈ H ₄ F ₄ O ₂ , F.W. 208.11 	1g 5g	115.00 455.00
H31863	5-Fluoro-2-(trifluoromethoxy)benzamide, JRD, 97% C ₈ H ₄ F ₄ NO ₂ , F.W. 223.12 	250mg 1g	67.70 226.00
H31623	5-Fluoro-2-(trifluoromethoxy)benzoic acid, JRD, 97% C ₈ H ₄ F ₄ O ₃ , F.W. 224.11 	1g 5g	114.00 457.00
H32804	5-Fluoro-2-(trifluoromethoxy)benzoyl chloride, JRD, 97%  C ₈ H ₃ ClF ₄ O ₂ , F.W. 242.55, UN3265 	250mg 1g	54.40 182.00
H32344	5-Fluoro-2-(trifluoromethoxy)cinnamic acid, JRD, 97% C ₁₀ H ₆ F ₄ O ₃ , F.W. 250.15 	250mg 1g	54.40 182.00
H32797	4-Fluoro-3-(trifluoromethoxy)phenylacetonitrile, 97% [1020718-23-3], C ₉ H ₅ F ₄ NO, F.W. 219.14, b.p. 91°/1mm, UN3276, MDL MFCD0772785 	250mg 1g	93.50 310.00

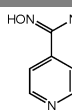
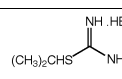
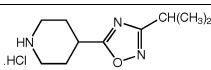
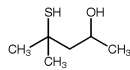
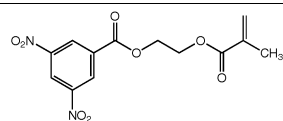
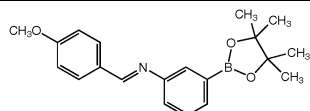
Stock #	Description		Size	\$
H32171	5'-Fluoro-2'-(trifluoromethyl)acetophenone, JRD, 97% C ₉ H ₆ F ₄ O, F.W. 206.04  R:36/38, S:26-37		1g 5g	74.40 298.00
H32684	3-Fluoro-2-(trifluoromethyl)aniline, JRD, 97% C ₈ H ₆ F ₃ N, F.W. 179.11, UN2810  R:20/21/22-36/38, S:26-36/37		250mg 1g	81.70 273.00
H32180	3-Fluoro-2-(trifluoromethyl)benzaldehyde, JRD, 97% Δ C ₈ H ₅ F ₃ O, F.W. 192.11  R:36/37/38, S:26-37		250mg 1g	40.80 136.00
H32210	3-Fluoro-2-(trifluoromethyl)benzylamine, JRD, 97% Δ [771581-62-5], C ₈ H ₇ F ₃ N, F.W. 193.14, UN2735  R:34, S:26-36/37/39-45		250mg 1g	81.70 273.00
H31916	3-Fluoro-2-(trifluoromethyl)phenylacetic acid, JRD, 97% [897940-14-6], C ₉ H ₆ F ₄ O ₂ , F.W. 222.14, m.p. 122-124°  R:36/37/38, S:26-37		250mg 1g	126.00 418.00
H31608	3-Fluoro-2-(trifluoromethyl)phenylacetonitrile, JRD, 97% C ₉ H ₆ F ₃ N, F.W. 203.14, UN3276  R:20/21/22-36/38, S:26-36/37		250mg 1g	109.00 365.00
H31874	3-Fluoro-2-(trifluoromethyl)-DL-phenylalanine, JRD, 97% C ₁₀ H ₉ F ₄ NO ₂ , F.W. 251.18		250mg	142.00
H31506	2-[2-Fluoro-3-(trifluoromethyl)phenyl]ethanol, JRD, tech. 90% C ₉ H ₈ F ₄ O, F.W. 208.15, n _D ²⁰ 1.4558		250mg 1g	33.60 112.00
H31530	2-[2-Fluoro-4-(trifluoromethyl)phenyl]ethanol, JRD, tech. 90% C ₉ H ₈ F ₄ O, F.W. 208.15, n _D ²⁰ 1.4552		250mg 1g	33.60 112.00
H32943	2-[2-Fluoro-5-(trifluoromethyl)phenyl]ethanol, JRD, 97% C ₉ H ₈ F ₄ O, F.W. 208.15		250mg 1g	33.60 112.00
H32867	2-[4-Fluoro-3-(trifluoromethyl)phenyl]ethanol, JRD, 97% C ₉ H ₈ F ₄ O, F.W. 208.15		250mg 1g	33.60 112.00
H32335	3-[2-Fluoro-3-(trifluoromethyl)phenyl]propionic acid, JRD, 97% C ₁₀ H ₈ F ₄ O ₂ , F.W. 236.16  R:36/37/38, S:26-37		250mg 1g	33.60 115.00
H31895	3-[2-Fluoro-4-(trifluoromethyl)phenyl]propionic acid, JRD, 97% C ₁₀ H ₈ F ₄ O ₂ , F.W. 236.16  R:36/37/38, S:26-37		250mg 1g	33.60 112.00
H32071	3-[2-Fluoro-5-(trifluoromethyl)phenyl]propionic acid, JRD, 97% C ₁₀ H ₈ F ₄ O ₂ , F.W. 236.16  R:36/37/38, S:26-37		250mg 1g	33.60 112.00
H32097	3-[3-Fluoro-4-(trifluoromethyl)phenyl]propionic acid, JRD, 97% C ₁₀ H ₈ F ₄ O ₂ , F.W. 236.16  R:36/37/38, S:26-37		250mg 1g	33.60 112.00
H32349	3-[4-Fluoro-2-(trifluoromethyl)phenyl]propionic acid, JRD, 97% C ₁₀ H ₈ F ₄ O ₂ , F.W. 236.16  R:36/37/38, S:26-37		250mg 1g	54.40 182.00

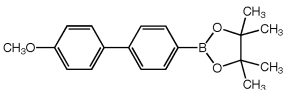
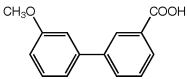
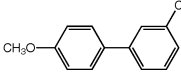
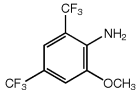
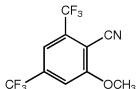
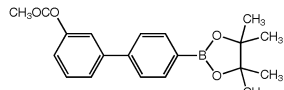
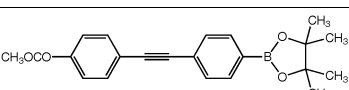
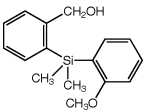
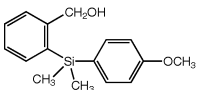
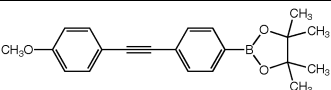
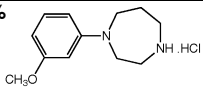
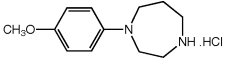
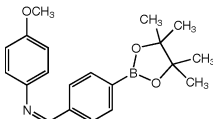
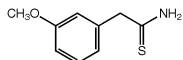
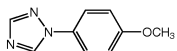
Stock #	Description		Size	\$
H31955	3-[4-Fluoro-3-(trifluoromethyl)phenyl]propionic acid, JRD, 97% C ₁₀ H ₈ F ₄ O ₂ , F.W. 236.16 ✗ R:36/37/38, S:26-37		250mg 1g	54.40 182.00
H32633	2-Fluoro-6-(trifluoromethyl)pyridine, 98% [94239-04-0], C ₆ H ₅ F ₄ N, F.W. 165.09, UN1993, EINECS 428-100-3 ✗ R:10-20/22-52/53, S:16-61		1g 5g	26.60 82.20
H51815	3-(Fmoc-amino)benzamidoxime, 97% C ₂₂ H ₁₉ N ₃ O ₃ , F.W. 373.41 ✗ R:36/37/38, S:26-37		250mg 1g	78.00 267.00
H51824	4-(Fmoc-amino)benzamidoxime, 97% C ₂₂ H ₁₉ N ₃ O ₃ , F.W. 373.41 ✗ R:36/37/38, S:26-37		250mg 1g	78.00 267.00
H51855	2-(Fmoc-amino)-4,5-diphenylthiophene-3-carboxylic acid, 97% C ₂₈ H ₂₃ NO ₄ S, F.W. 397.49 ✗ R:36/37/38, S:26-37		250mg 1g	55.00 165.00
H51843	2-(Fmoc-amino)-5-methyl-4-phenylthiophene-3-carboxylic acid, 97% C ₂₇ H ₂₁ NO ₄ S, F.W. 455.53 ✗ R:36/37/38, S:26-37		250mg 1g	44.00 133.00
H51721	6-(Fmoc-amino)nicotinamidoxime, 97% C ₂₁ H ₁₈ N ₄ O ₃ , F.W. 374.40, m.p. 158-162° ✗ R:36/37/38, S:26-37		250mg 1g	132.00 377.00
H51802	2-(Fmoc-amino)-4-phenylthiophene-3-carboxylic acid, 97% C ₂₆ H ₁₉ NO ₄ S, F.W. 441.50 ✗ R:36/37/38, S:26-37		250mg 1g	42.40 127.00
H51823	3-(Fmoc-amino)thiobenzamide, 97% C ₂₂ H ₁₈ N ₂ O ₂ S, F.W. 374.46 ✗ R:36/37/38, S:26-37		250mg 1g	44.00 152.00
H51829	4-(Fmoc-amino)thiobenzamide, 97% C ₂₂ H ₁₈ N ₂ O ₂ S, F.W. 374.46 ✗ R:36/37/38, S:26-37		250mg 1g	44.00 152.00
H31053	N-Formylglycine, 98% [For-Gly-OH] [2491-15-8], HCONHCH ₂ CO ₂ H, F.W. 103.08, m.p. 147-151°, EINECS 219-649-8, RTECS MC0547000		1g 5g	29.80 103.00
H51755	2-[(4-Formylphenyl)dimethylsilyl]benzyl alcohol, 95% C ₁₈ H ₁₈ O ₂ Si, F.W. 270.40, Note: Product of Advanced Molecular Technologies Pty Ltd. ✗ R:36/37/38, S:26-37		1g 5g	67.50 270.00
H32013	2-Furaldehyde 2,2-dimethylhydrazone, 98% [14064-21-2], C ₇ H ₁₀ N ₂ O, F.W. 138.17, b.p. 69°/2mm, f.p. 92°(198°F), d. 1.048, n _D ²⁰ 1.5800 ✗ R:22-36/37/38, S:23-26-36/37		10g 50g	47.40 162.00
H51831	Furan-2-thiocarboxamide, 97% [17572-09-7], C ₅ H ₆ NOS, F.W. 127.16 ✗ R:36/37/38, S:26-37		250mg 1g	92.00 315.00
H51801	2-(2-Furanyl)-4-methylthiazole-5-carboxylic acid, 97% C ₉ H ₇ NO ₃ S, F.W. 209.22 ✗ R:36/37/38, S:26-37		250mg 1g	128.00 384.00
H51121	4-(Furfuryliminomethyl)benzeneboronic acid pinacol ester, 97% C ₁₈ H ₂₂ BNO ₃ , F.W. 311.18, MDL MFCD11113038		1g 5g 25g	29.20 112.00 453.00

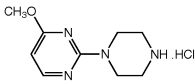
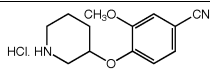
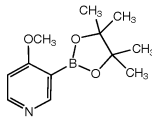
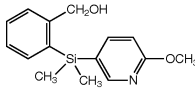
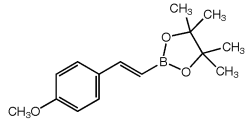
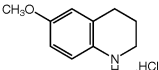
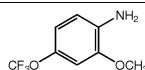
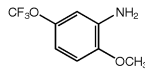
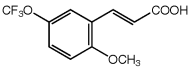
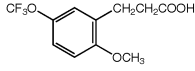
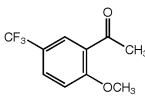
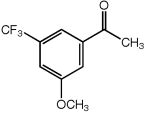
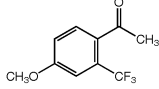
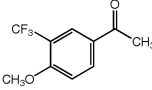
Stock #	Description		Size	\$
H32966	Furo[3,2-b]pyridine, 98% [272-62-8], C ₇ H ₅ NO, F.W. 119.12, b.p. 79-80°/12mm, n _D ²⁰ 1.5753  R:36/38, S:26-37		250mg	70.00
			1g	195.00
			5g	650.00
H30686	2-Furyldimethylsilanol sodium salt [879904-88-8], C ₆ H ₇ NaO ₂ Si, F.W. 164.21, m.p. ca 92° dec., MDL MFCD08457663  R:36/37/38, S:26-37		250mg	74.50
			1g	206.00
			5g	626.00
45580	Gallium(III) perchlorate hydrate, 13.5-15.5% Ga [81029-07-4], Ga(ClO ₄) ₃ ·xH ₂ O, F.W. 368.07(anhy), m.p. 175°, UN1481  R:8-36/37/38, S:17-26-37		2g	17.00
			10g	48.00
45572	Gallium(III) selenide, 99.99% (metals basis) [12024-24-7], Ga ₂ Se ₃ , F.W. 376.33, UN3077, MDL MFCD00016103, †  R:23/25-33-50/53, S:20/21-28-45-60-61		1g	27.80
			5g	125.00
45569	Gibberellic acid, 98% [77-06-5], C ₁₉ H ₂₂ O ₆ , F.W. 346.37, m.p. ca 230° dec., Merck 14,4419, EINECS 201-001-0, RTECS LY8990000, BRN 54346, MDL MFCD00079329, †		500mg	27.80
			2g	56.80
			10g	226.00
45649	Guanosine, 97% [118-00-3], C ₁₀ H ₁₃ N ₅ O ₆ , F.W. 283.24, m.p. ca 250° dec., Merck 14,4566, EINECS 204-227-8, RTECS MF8750000, BRN 625911, MDL MFCD00010182, †		100g	68.00
			500g	275.00
45612	Hafnium(IV) chloride, 99.9% (metals basis), Zr<0.5%  [13499-05-3], HfCl ₄ , F.W. 320.3, m.p. 320° subl., Merck 14,4588, UN3260, EINECS 236-826-5, MDL MFCD00011034, †  R:34, S:26-36/37/39-45		25g	46.30
			100g	167.00
			500g	750.00
45483	Hafnium(IV) oxide, 99.995%, (metals basis excluding Zr), Zr typically <0.2% [12055-23-1], HfO ₂ , F.W. 210.49, m.p. 2774°, d. 9.68, Merck 14,4588, EINECS 235-013-2, MDL MFCD00003565, †		1g	124.00
			5g	468.00
H51173	1-Heptylmagnesium chloride, 1M in MeTHF  [42092-01-3], CH ₃ (CH ₂) ₆ CH ₂ MgCl, F.W. 158.95, UN2924 R:11-14-19-34, S:8-16-26-36/37/39-43-45		100ml	47.30
			500ml	178.00
H51744	4-(Hexamethyleneiminomethyl)benzeneboronic acid pinacol ester, 95% C ₁₈ H ₃₀ BNO ₂ , F.W. 315.26  R:36/37/38, S:26-37		1g	76.90
			5g	310.00
45652	Hexanes, mixed isomers, HPLC Grade, 99+% [92112-69-1], C ₆ H ₁₄ , F.W. 86.18, m.p. -95°, b.p. 65-70°, f.p. <-23°(-9°F), d. 0.67, n _D ²⁰ 1.3795, UN1208, EINECS 295-570-2, †  R:11-38-48/20-62-51/53-65-67, S:9-16-29-33-36/37-61-62		1L	32.60
			4L	97.50
			4x1L	112.00
H51167	1-Hexylmagnesium chloride, 1M in MeTHF  [44767-62-6], CH ₃ (CH ₂) ₅ CH ₂ MgCl, F.W. 144.93, UN2924 R:11-14-19-34, S:8-16-26-36/37/39-43-45		100ml	40.70
			500ml	153.00
H30426	3-Hydroxybutyric acid, 98%  [300-85-6], C ₄ H ₈ O ₃ , F.W. 104.10, m.p. 36-39°, EINECS 206-099-9, †  R:36/37/38, S:26-37		5g	133.00
			25g	400.00
H51126	3-(Hydroxymethyl)benzeneboronic acid pinacol ester [443776-76-9], C ₁₃ H ₁₉ BO ₃ , F.W. 234.10, MDL MFCD09266196		1g	26.70
			5g	102.00
			25g	414.00
H51123	4-(Hydroxymethyl)benzeneboronic acid pinacol ester [302348-51-2], C ₁₃ H ₁₉ BO ₃ , F.W. 234.10, MDL MFCD09837617		1g	26.70
			5g	103.00
			25g	414.00

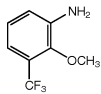
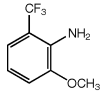
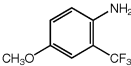
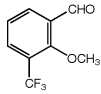
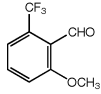
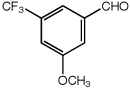
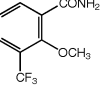
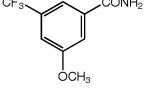
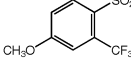
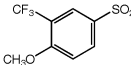
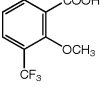
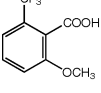
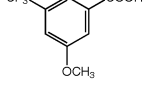
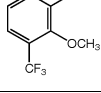
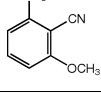
Stock #	Description		Size	\$
H30624	3'-Hydroxy-6-methylflavone, 97% C ₁₆ H ₁₂ O ₃ , F.W. 252.28, m.p. 203-206°, MDL MFCD07778551 ✗ R:36/37/38, S:26-37		1g	33.50
			5g	103.00
			25g	316.00
H30959	4'-Hydroxy-6-methylflavone, 97% C ₁₆ H ₁₂ O ₃ , F.W. 252.28, m.p. 238-243°, MDL MFCD03424432 ✗ R:36/37/38, S:26-37		1g	36.60
			5g	112.00
			25g	345.00
H51880	2-Hydroxy-3-methylpyrazine, 96% [19838-07-4], C ₅ H ₆ N ₂ O, F.W. 110.11, m.p. 152-153°, EINECS 243-362-7 ✗ R:36/37/38, S:26-37		250mg	95.00
			1g	306.00
H51796	2-Hydroxy-6-methylpyridine-3-carboxamidoxime, 97% C ₇ H ₈ N ₃ O ₂ , F.W. 167.17 ✗ R:36/37/38, S:26-37		250mg	92.00
			1g	315.00
H32921	4-(Hydroxymethyl)pyridine-2-carboxylic acid, 95% [923169-37-3], C ₇ H ₇ NO ₃ , F.W. 153.14 ✗ R:36/37/38, S:26-37		1g	34.80
			5g	116.00
			25g	385.00
H30725	5-Hydroxynicotinic acid, 97% [5-Hydroxypyridine-3-carboxylic acid] [27828-71-3], C ₆ H ₅ NO ₃ , F.W. 139.11, m.p. >300°, RTECS QT1757500 ✗ R:36/37/38, S:26-37		1g	37.50
			5g	124.00
			25g	414.00
H51726	trans-4-Hydroxy-L-proline methyl ester hydrochloride, 98% [40216-83-9], C ₆ H ₁₁ NO ₃ ·HCl, F.W. 181.62, BRN 4716932, MDL MFCD00080855		5g	31.10
			25g	120.00
H51746	4-(3-Hydroxypropylcarbamoyl)benzeneboronic acid pinacol ester, 95% C ₁₆ H ₂₄ BNO ₄ , F.W. 305.18		1g	54.40
			5g	220.00
H31133	(2-Hydroxypropyl)-β-cyclodextrin, MW ca 1250-1480 [128446-35-5], m.p. 305°, EINECS 420-920-1, RTECS GU2293332, MDL MFCD00069372		5g	36.10
			25g	118.00
			100g	328.00
H51901	4-(3-Hydroxy-1-propynyl)benzeneboronic acid pinacol ester, 95% C ₁₅ H ₁₉ BO ₃ , F.W. 258.12		1g	60.00
			5g	240.00
H51673	6-Hydroxypyridine-3-boronic acid pinacol ester, 97% C ₁₁ H ₁₆ BNO ₃ , F.W. 221.06		250mg	64.50
			1g	205.00
H51096	5-Hydroxyquinoline, 99% [578-67-6], C ₉ H ₇ NO, F.W. 145.16, EINECS 209-428-4, MDL MFCD00006792 ✗ R:36/37/38, S:26-37		250mg	49.70
			1g	141.00
H32780	4-Hydroxythiobenzamide, 98% [25984-63-8], C ₇ H ₇ NOS, F.W. 153.20, m.p. 181-185° ✗ R:22-36/37/38, S:26-36/37		5g	94.20
			25g	313.00
H31734	5-Hydroxy-2-(trifluoromethyl)pyridine, 97% [216766-12-0], C ₆ H ₄ F ₃ NO, F.W. 163.10, m.p. 174-176°, MDL MFCD06801325 ✗ R:36/37/38, S:26-37		250mg	64.60
			1g	179.00

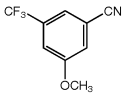
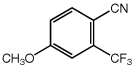
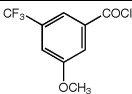
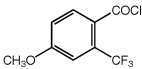
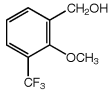
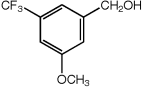
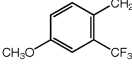
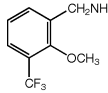
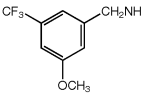
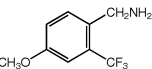
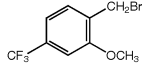
Stock #	Description		Size	\$
H32992	(4-Imidazolyl)acetonitrile, 97% [18502-05-1], C ₅ H ₅ N ₃ , F.W. 107.11, m.p. 136-138°, UN3439, MDL MFCD00070306  R:25-36/37/38, S:26-36/37-45		1g	107.00
			5g	320.00
H51671	4-[3-(1-Imidazolyl)propylaminomethyl]benzeneboronic acid pinacol ester C ₁₉ H ₂₈ BN ₂ O ₅ , F.W. 341.26  R:36/37/38, S:26-37		1g	47.50
			5g	190.00
45525	Indium(III) fluoride, anhydrous, 99.99% (metals basis) ▴ [7783-52-0], InF ₃ , F.W. 171.82, m.p. ca 1170°, b.p. >1200°, d. 4.39, Merck 14,4959, UN2923, EINECS 232-005-0, MDL MFCD00016149, †  R:23/25-34, S:9-20-26-36/37/39-45		5g	62.40
			25g	280.00
45563	Indium(III) sulfide, 99.999% (metals basis) [12030-24-9], In ₂ S ₃ , F.W. 325.83, m.p. 1050°, EINECS 234-742-3, MDL MFCD00011061, †  R:20/22-32-36/37/38, S:7-26-36		2g	51.30
			10g	144.00
			50g	650.00
H51075	(R)-(-)-4-(3-Indolylmethyl)-2-oxazolidinone, 98% ▴ [157636-81-2], C ₁₂ H ₁₂ N ₂ O ₃ , F.W. 216.24, MDL MFCD01863583  R:36/37/38, S:26-37		1g	178.00
H51119	4-Iodobenzeneboronic acid ▴ [5122-99-6], C ₆ H ₅ BiO ₂ , F.W. 247.83, MDL MFCD02093073  R:22-36/37/38, S:26-36/37		1g	28.40
			25g	439.00
H31826	4-Iodobutyl acetate, 97%, stab. with copper ▴ [40596-44-9], CH ₃ CO ₂ (CH ₂) ₄ I, F.W. 242.05, b.p. 94-95°/5.5mm, f.p. 91°(196°F), d. 1.610, n _D ²⁰ 1.5500		10g	61.10
			50g	191.00
H51875	2-Iodo-4,6-dimethylaniline, 98% ▴ [4102-54-9], C ₈ H ₁₀ IN, F.W. 247.08, m.p. 64-65°, UN2811  R:20/21/22-36/38, S:26-36/37		1g	45.00
			5g	191.00
H51868	2-Iodoimidazole, 96% ▴ [3034-62-6], C ₃ H ₃ IN ₂ , F.W. 193.97, m.p. 190-191°, MDL MFCD00159701  R:36/37/38, S:26-37		250mg	69.00
			1g	192.00
H51862	2-Iodo-1-methylimidazole, 97% ▴ [37067-95-1], C ₄ H ₅ IN ₂ , F.W. 208.00, m.p. 79-82°, MDL MFCD00661606  R:36/37/38, S:26-37		250mg	64.00
			1g	188.00
H30803	3-Iodophenyl isocyanate, 97% ▴ ▴ [23138-56-9], C ₇ H ₄ INO, F.W. 245.02, b.p. 55°/0.2mm, d. 1.860, UN2206  R:20/22-36/37/38-42, S:23-26-36/37-45		1g	59.20
			5g	243.00
H31468	2-Iodopyridine-4-carboxamide, 95% ▴ [2-Iodoisonicotinamide] C ₆ H ₅ IN ₂ O, F.W. 248.02, m.p. 164-166°  R:36/37/38, S:26-37		250mg	62.20
			1g	173.00
			5g	570.00
H31002	2-Iodopyridine-4-carboxylic acid, 95% ▴ [2-Iodoisonicotinic acid] [58481-10-0], C ₆ H ₄ INO ₂ , F.W. 249.01, m.p. ca 210° dec.  R:36/37/38, S:26-37		250mg	78.30
			1g	206.00
H51871	2-Iodo-1-tritylimidazole, 97% ▴ [67478-46-0], C ₂₂ H ₁₇ IN ₂ , F.W. 436.29, m.p. 170-171°  R:36/37/38, S:26-37		250mg	51.00
			1g	150.00
45570	Iridium, 1% on 5mm alumina spheres, reduced, anhydrous MDL MFCD00011062, †		10g	99.40
			50g	333.00
			250g	1303.00
H32278	Iron(III) 1,3-diphenyl-1,3-propanedionate C ₁₅ H ₁₃ FeO ₆ , F.W. 725.61		5g	77.50
			25g	232.00

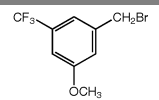
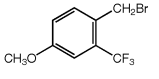
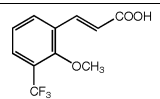
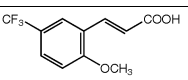
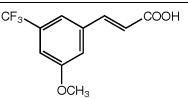
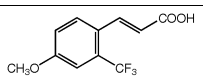
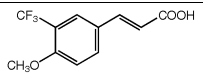
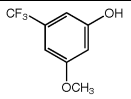
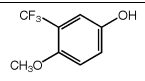
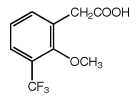
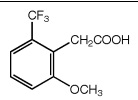
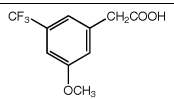
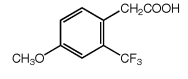
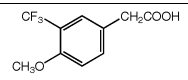
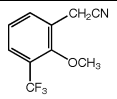
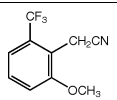
Stock #	Description		Size	\$
H51722	Isonicotinamidoxime, 97% [1594-57-6], C ₆ H ₇ N ₃ O, F.W. 137.14, m.p. 212-216°, RTECS NS1050000, MDL MFCD00125873 ☒ R:36/37/38, S:26-37		1g 5g	57.90 200.00
H51852	S-Isopropylisothiurea hydrobromide, 97% [57200-31-4], C ₄ H ₁₀ N ₂ S HBr, F.W. 199.11 ☒ R:36/37/38, S:26-37		1g 5g 25g	24.00 107.00 441.00
H51155	Isopropylmagnesium chloride, 1M in MeTHF △ ▽ [1068-55-9], (CH ₃) ₂ CHMgCl, F.W. 102.85, UN3399, † ☒ R:11-14/15-19-34, S:8-16-26-36/37/39-43-45		100ml 500ml	28.10 70.10
H51156	Isopropylmagnesium chloride - LiCl complex, 1M in MeTHF △ ▽ [807329-97-1], (CH ₃) ₂ CHMgCl·LiCl, F.W. 145.24, f.p. -17°(1.4°F), d. 0.951, UN3399 ☒ R:11-14/15-19-34, S:8-16-26-36/37/39-43-45		100ml 500ml	78.80 328.00
H51025	4-(3-Isopropyl-1,2,4-oxadiazol-5-yl)piperidine hydrochloride C ₁₀ H ₁₇ N ₃ O HCl, F.W. 231.73, MDL MFCD11111021 ☒ R:36/37/38, S:26-37		250mg 1g	60.30 216.00
45501	Lithium aluminum hydride, 4% in THF △ ▽ [16853-85-3], LiAlH ₄ , F.W. 37.95, d. 0.90, Merck 14,349, Application(s): Reducing agent for pharmaceuticals and organic chemicals, UN3399, EINECS 240-877-9, MDL MFCD00011075, † ☒ R:11-15-19-34, S:7/8-16-26-36/37/39-43-45		500ml	134.00
45502	Lithium aluminum hydride, 10% in THF △ ▽ [16853-85-3], LiAlH ₄ , F.W. 37.95, d. 0.90, Merck 14,349, Application(s): Reducing agent for pharmaceuticals and organic chemicals, UN3399, EINECS 240-877-9, MDL MFCD00011075, † ☒ R:11-15-19-34, S:7/8-16-26-36/37/39-43-45		500ml	230.00
45503	Lithium aluminum hydride, 15% in THF/toluene (2.4:1) △ ▽ [16853-85-3], LiAlH ₄ , F.W. 37.95, d. 0.90, Merck 14,349, Application(s): Reducing agent for pharmaceuticals and organic chemicals, UN3399, EINECS 240-877-9, MDL MFCD00011075, † ☒ R:11-15-19-34-48/20-63-67, S:7/8-16-26-36/37/39-43-45		500ml	391.00
45491	Lithium dimethylamide, 10% w/v in hexane △ ▽ [3585-33-9], LiN(CH ₃) ₂ , F.W. 51.02, UN3394, EINECS 222-714-3, MDL MFCD00008295 ☒ R:14-17-34-48/20-62-51/53-65-67, S:9-16-26-36/37/39-45-61-62		50ml 250ml	42.80 171.00
46120	Magnesium bromide hexahydrate, 98+% ■ [13446-53-2], MgBr ₂ ·6H ₂ O, F.W. 292.22 (184.13anhy), m.p. 172°, d. 2.00, EINECS 232-170-9, MDL MFCD00149780, † ☒ R:36/37/38, S:26		100g 500g 2kg	21.10 78.50 255.00
45517	Magnesium germanide, 99.99% (metals basis) [1310-52-7], Mg ₂ Ge, F.W. 121.22, MDL MFCD11044441, †		5g 25g	158.00 640.00
45518	Magnesium silicide, 99.99% (metals basis) [22831-39-6], Mg ₂ Si, F.W. 76.71, Merck 14,5688, UN2624, EINECS 245-254-5, MDL MFCD00016202, † ☒ R:15, S:7/8-33-43		10g 50g 250g	32.10 118.00 521.00
46114	Magnesium thiosulfate hexahydrate, 99% [Magnesium hyposulfite] [10124-53-5], MgS ₂ O ₃ ·6H ₂ O, F.W. 244.53 (136.44anhy), m.p. 1700° dec., d. 1.818, Merck 14,5694, Solubility: Soluble in water. Insoluble in alcohol, EINECS 233-340-5, MDL MFCD00169871, † ☒ R:36/37/38, S:26		100g 500g	30.80 93.20
45557	Mercaptoacetic acid calcium salt trihydrate, 98% [Calcium mercaptoacetate trihydrate, Calcium thioglycolate trihydrate] [65208-41-5], (HSCH ₂ CO ₂)Ca·3H ₂ O, F.W. 276.34 (222.29anhy), m.p. 280-290° dec., Merck 14,1711, EINECS 212-402-5, BRN 3755791, MDL MFCD00150707, † ☒ R:22, S:36		10g 50g	23.30 87.50
H32107	4-Mercapto-4-methyl-2-pentanol, 98% [31539-84-1], C ₆ H ₁₄ OS, F.W. 134.24, b.p. 40°/0.6mm, f.p. 70°(158°F), d. 0.957, n _D ²⁰ 1.4710 ☒ R:22-36/37/38, S:26-36/37		250mg 1g	89.30 250.00
H30745	2-(Methacryloyloxy)ethyl 3,5-dinitrobenzoate, 97% C ₁₃ H ₁₂ N ₂ O ₈ , F.W. 324.25, m.p. 70-72° ☒ R:36/37/38, S:26-28		5g 25g	55.30 184.00
H51129	3-(4-Methoxybenzylideneamino)benzeneboronic acid pinacol ester [380151-91-7], C ₂₀ H ₂₄ BNO ₃ , F.W. 337.22, MDL MFCD11113043		1g 5g 25g	24.20 92.60 375.00

Stock #	Description		Size	\$
H51907	4'-Methoxybiphenyl-4-boronic acid pinacol ester, 95% C ₁₈ H ₂₃ BO ₃ , F.W. 310.20		1g	51.60
			5g	210.00
H51767	3'-Methoxybiphenyl-3-carboxylic acid, 95% [168618-45-9], C ₁₄ H ₁₂ O ₃ , F.W. 228.25 ✖ R:36/37/38, S:26-37		1g	60.80
			5g	195.00
H51764	4'-Methoxybiphenyl-3-carboxylic acid, 95% [725-05-3], C ₁₄ H ₁₂ O ₃ , F.W. 228.25 ✖ R:36/37/38, S:26-37		1g	60.80
			5g	160.00
H31582	2-Methoxy-4,6-bis(trifluoromethyl)aniline, JRD, 97% C ₈ H ₇ F ₃ NO, F.W. 259.15, UN2811 ✖ R:20/21/22-36/38, S:26-36/37		250mg	54.40
			1g	185.00
H31652	2-Methoxy-4,6-bis(trifluoromethyl)benzonitrile, JRD, 97% C ₁₀ H ₅ F ₃ NO, F.W. 269.14, m.p. 46-48°, UN3439 ✖ R:20/21/22-36/38, S:26-36/37		250mg	54.40
			1g	182.00
H51916	3'-(Methoxycarbonyl)biphenyl-4-boronic acid pinacol ester, 95% C ₂₀ H ₂₃ BO ₄ , F.W. 338.21		1g	62.50
			5g	250.00
H51900	4-[4-(Methoxycarbonyl)phenylethynyl]-benzeneboronic acid pinacol ester, 95% C ₂₂ H ₂₃ BO ₄ , F.W. 362.23		1g	65.60
			5g	260.00
H51742	2-[(2-Methoxyphenyl)dimethylsilyl]benzyl alcohol, 95% C ₁₆ H ₂₀ O ₂ Si, F.W. 272.41, Note: Product of Advanced Molecular Technologies Pty Ltd. ✖ R:36/37/38, S:26-37		1g	48.40
			5g	190.00
H51662	2-[(4-Methoxyphenyl)dimethylsilyl]benzyl alcohol [944064-51-1], C ₁₆ H ₂₀ O ₂ Si, F.W. 272.41, MDL MFCD11870726, Note: Product of Advanced Molecular Technologies Pty Ltd. ✖ R:36/37/38, S:26-37		1g	41.60
			5g	156.00
H51914	4-(4-Methoxyphenylethynyl)benzeneboronic acid pinacol ester, 95% C ₂₁ H ₂₃ BO ₃ , F.W. 334.22		1g	51.60
			5g	210.00
H51751	1-(3-Methoxyphenyl)homopiperazine monohydrochloride, 98% C ₁₂ H ₁₈ N ₂ O·HCl, F.W. 242.74, m.p. 169-173°, Note: Sold in collaboration with Tosoh Organic Chemical Co., Ltd.		500mg	390.00
H51703	1-(4-Methoxyphenyl)homopiperazine monohydrochloride, 98% [934992-02-6], C ₁₂ H ₁₈ N ₂ O·HCl, F.W. 242.74, m.p. 175-179°, Note: Sold in collaboration with Tosoh Organic Chemical Co., Ltd. ✖ R:36/37/38, S:26-37		500mg	390.00
H51116	4-(4-Methoxyphenyliminomethyl)benzeneboronic acid pinacol ester C ₂₀ H ₂₄ BNO ₃ , F.W. 337.22, MDL MFCD10567027		1g	24.20
			5g	92.60
			25g	375.00
H31615	2-(3-Methoxyphenyl)thioacetamide, 97% [35582-11-7], C ₉ H ₁₁ NOS, F.W. 181.26, m.p. 104-108° ✖ R:22-36/37/38, S:26-36/37		1g	97.00
			5g	325.00
H30962	1-(4-Methoxyphenyl)-1H-1,2,4-triazole, 99% [68377-33-3], C ₈ H ₈ N ₃ O, F.W. 175.19, m.p. 96-98 ✖ R:36/37/38, S:26-37		5g	58.90
			25g	196.00

Stock #	Description		Size	\$
H51678	4-Methoxy-2-(1-piperazinyl)pyrimidine hydrochloride, 96% C ₉ H ₁₄ N ₄ O·HCl, F.W. 230.71		250mg 1g	59.70 199.00
H51138	3-Methoxypiperidine hydrochloride ■ [688809-94-1], C ₆ H ₁₃ NO·HCl, F.W. 151.64, MDL MFCD06800959 ✗ R:36/37/38, S:26-37		250mg 1g	61.00 219.00
H51677	3-Methoxy-4-(3-piperidinyloxy)benzonitrile hydrochloride, 97% C ₁₃ H ₁₆ N ₂ O ₂ ·HCl, F.W. 268.7 ✗ R:22-36/37/38, S:26-36/37		250mg 1g	61.30 219.00
H30555	4-Methoxypyridine-3-boronic acid pinacol ester, 97% [758699-74-0], C ₁₂ H ₁₅ BO ₃ , F.W. 235.09		1g 5g	114.00 380.00
H51740	2-[(6-Methoxy-3-pyridyl)dimethylsilyl]benzyl alcohol, 95% C ₁₉ H ₁₉ NO ₂ Si, F.W. 273.40, Note: Product of Advanced Molecular Technologies Pty Ltd. ✗ R:36/37/38, S:26-37		1g 5g	62.00 260.00
H30373	4-Methoxy-trans-β-styrylboronic acid pinacol ester, 98% [2-[trans-2-(4-Methoxyphenyl)vinyl]-4,4,5,5-tetramethyl-1,3,2]dioxaborolane] [149777-83-3], C ₁₅ H ₂₁ BO ₃ , F.W. 260.14		250mg 1g 5g	47.40 132.00 441.00
H31910	6-Methoxy-1,2,3,4-tetrahydroisoquinoline hydrochloride, 95% [57196-62-0], C ₁₀ H ₁₃ NO·HCl, F.W. 199.68, m.p. 233-236°, MDL MFCD01717328 ✗ R:36/37/38, S:26-37		250mg 1g	98.90 275.00
H32743	2-Methoxy-4-(trifluoromethoxy)aniline, JRD, 97% C ₈ H ₈ F ₃ NO ₂ , F.W. 207.15, m.p. 37-39°, UN2811 ✗ R:20/21/22-33-36/38, S:26-28-36/37		1g 5g	117.00 468.00
H32219	2-Methoxy-5-(trifluoromethoxy)aniline, JRD, 97% C ₈ H ₈ F ₃ NO ₂ , F.W. 207.15, UN2810 ✗ R:20/21/22-33-36/38, S:26-28-36/37		1g 5g	117.00 467.00
H32052	2-Methoxy-5-(trifluoromethoxy)cinnamic acid, JRD, 97% C ₁₁ H ₉ F ₃ O ₄ , F.W. 262.18 ✗ R:36/37/38, S:26-37		250mg 1g	33.60 112.00
H32321	3-[2-Methoxy-5-(trifluoromethoxy)phenyl]propionic acid, JRD, 97% C ₁₁ H ₁₁ F ₃ O ₄ , F.W. 264.20 ✗ R:36/37/38, S:26-37		250mg 1g	67.70 226.00
H32652	2'-Methoxy-5'-(trifluoromethyl)acetophenone, JRD, 97% [503464-99-1], C ₁₀ H ₉ F ₃ O ₂ , F.W. 218.17 ✗ R:36/37/38, S:26-37		1g 5g	105.00 420.00
H32322	3'-Methoxy-5'-(trifluoromethyl)acetophenone, JRD, 97% C ₁₀ H ₉ F ₃ O ₂ , F.W. 218.17 ✗ R:36/37/38, S:26-37		250mg 1g	54.40 182.00
H31627	4'-Methoxy-2'-(trifluoromethyl)acetophenone, JRD, 97% C ₁₀ H ₉ F ₃ O ₂ , F.W. 218.17 ✗ R:36/37/38, S:26-37		250mg 1g	54.40 182.00
H32809	4'-Methoxy-3'-(trifluoromethyl)acetophenone, JRD, 97% C ₁₀ H ₉ F ₃ O ₂ , F.W. 218.17 ✗ R:36/37/38, S:26-37		1g 5g	56.40 225.00

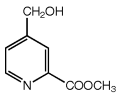
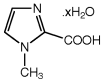
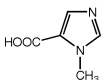
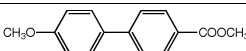
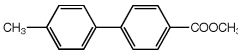
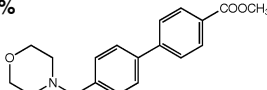
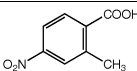
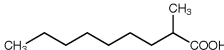
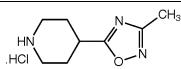
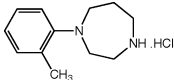
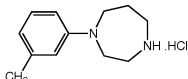
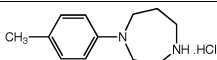
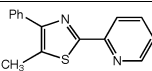
Stock #	Description		Size	\$
H32045	2-Methoxy-3-(trifluoromethyl)aniline, JRD, 97% C ₈ H ₆ F ₃ NO, F.W. 191.15, UN2810 ☒ R:20/21/22-33-36/38, S:26-28-36/37		1g 5g	117.00 467.00
H32449	2-Methoxy-6-(trifluoromethyl)aniline, JRD, 97% C ₈ H ₆ F ₃ NO, F.W. 191.15, UN2810 ☒ R:20/21/22-33-36/38, S:26-28-36/37		1g 5g	117.00 470.00
H31872	4-Methoxy-2-(trifluoromethyl)aniline, JRD, 97% C ₈ H ₆ F ₃ NO, F.W. 191.15, UN2810 ☒ R:20/21/22-33-36/38, S:26-28-36/37		250mg 1g	54.40 182.00
H31642	2-Methoxy-3-(trifluoromethyl)benzaldehyde, JRD, 97% △ C ₈ H ₅ F ₃ O ₂ , F.W. 204.15 ☒ R:36/37/38, S:26-37		1g 5g	88.60 355.00
H32206	2-Methoxy-6-(trifluoromethyl)benzaldehyde, JRD, 97% △ C ₈ H ₅ F ₃ O ₂ , F.W. 204.15, m.p. 100-103° ☒ R:36/37/38, S:26-37		1g 5g	103.00 411.00
H32791	3-Methoxy-5-(trifluoromethyl)benzaldehyde, JRD, 97% △ C ₈ H ₅ F ₃ O ₂ , F.W. 204.15 ☒ R:36/37/38, S:26-37		1g 5g	68.10 273.00
H32690	2-Methoxy-3-(trifluoromethyl)benzamide, JRD, 97% C ₈ H ₆ F ₃ NO ₂ , F.W. 219.16 ☒ R:36/37/38, S:26-37		1g 5g	105.00 415.00
H31654	3-Methoxy-5-(trifluoromethyl)benzamide, JRD, 97% C ₈ H ₆ F ₃ NO ₂ , F.W. 219.16 ☒ R:36/37/38, S:26-37		1g 5g	103.00 411.00
H32280	4-Methoxy-2-(trifluoromethyl)benzenesulfonyl chloride, JRD, 97% ▴ C ₈ H ₆ ClF ₃ O ₃ S, F.W. 274.64, m.p. 57-60°, UN3261 ☒ R:34, S:26-36/37/39-45		250mg 1g	67.70 226.00
H32242	4-Methoxy-3-(trifluoromethyl)benzenesulfonyl chloride, JRD, 97% ▴ C ₈ H ₆ ClF ₃ O ₃ S, F.W. 274.64, m.p. 52-55°, UN3261 ☒ R:34, S:26-36/37/39-45		250mg 1g	67.70 230.00
H32198	2-Methoxy-3-(trifluoromethyl)benzoic acid, JRD, 97% C ₈ H ₅ F ₃ O ₃ , F.W. 220.15 ☒ R:36/37/38, S:26-37		1g 5g	88.60 355.00
H32789	2-Methoxy-6-(trifluoromethyl)benzoic acid, JRD, 97% [119692-41-0], C ₈ H ₅ F ₃ O ₃ , F.W. 220.15 ☒ R:36/37/38, S:26-37		1g 5g	105.00 415.00
H32430	3-Methoxy-5-(trifluoromethyl)benzoic acid, JRD, 97% C ₈ H ₅ F ₃ O ₃ , F.W. 220.15 ☒ R:36/37/38, S:26-37		1g 5g	69.70 280.00
H32611	2-Methoxy-3-(trifluoromethyl)benzonitrile, JRD, 97% C ₈ H ₅ F ₃ NO, F.W. 201.15, UN3276 ☒ R:20/21/22-36/38, S:26-36/37		1g 5g	88.60 355.00
H32470	2-Methoxy-6-(trifluoromethyl)benzonitrile, JRD, 97% C ₈ H ₅ F ₃ NO, F.W. 201.15, m.p. 66-68°, UN3439 ☒ R:20/21/22-36/38, S:26-36/37		1g	117.00

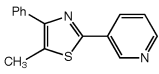
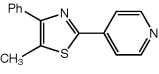
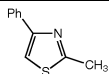
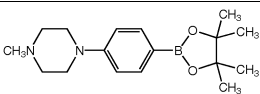
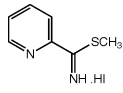
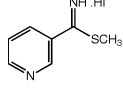
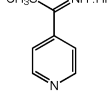
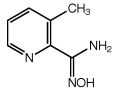
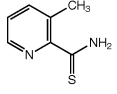
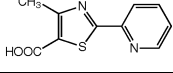
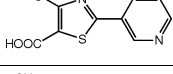
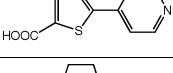
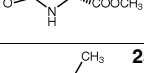
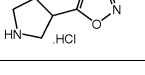
Stock #	Description		Size	\$
H32508	3-Methoxy-5-(trifluoromethyl)benzonitrile, JRD, 97% C ₉ H ₆ F ₃ NO, F.W. 201.15, UN3439  R:20/21/22-36/37/38, S:26-36/37		1g	124.00
			5g	495.00
H31513	4-Methoxy-2-(trifluoromethyl)benzonitrile, JRD, 97% C ₉ H ₆ F ₃ NO, F.W. 201.15, m.p. 63-66°, UN3439  R:20/21/22-36/38, S:26-36/37		1g	88.60
			5g	355.00
H32311	2-Methoxy-6-(trifluoromethyl)benzoyl chloride, JRD, 97%  C ₉ H ₆ ClF ₃ O ₂ , F.W. 238.59, UN3261  R:34, S:26-36/37/39-45		1g	117.00
			5g	467.00
H32703	3-Methoxy-5-(trifluoromethyl)benzoyl chloride, JRD, 97%  C ₉ H ₆ ClF ₃ O ₂ , F.W. 238.59, UN3265  R:34, S:26-36/37/39-45		1g	88.60
			5g	355.00
H31821	4-Methoxy-2-(trifluoromethyl)benzoyl chloride, JRD, 97%  C ₉ H ₆ ClF ₃ O ₂ , F.W. 238.59, UN3265  R:34, S:26-36/37/39-45		1g	56.40
			5g	225.00
H32404	2-Methoxy-3-(trifluoromethyl)benzyl alcohol, JRD, 97% C ₉ H ₈ F ₃ O ₂ , F.W. 206.16  R:36/37/38, S:26-37		1g	117.00
			5g	470.00
H32539	2-Methoxy-6-(trifluoromethyl)benzyl alcohol, JRD, 97% C ₉ H ₈ F ₃ O ₂ , F.W. 206.16  R:36/37/38, S:26-37		250mg	40.80
			1g	136.00
H32555	3-Methoxy-5-(trifluoromethyl)benzyl alcohol, JRD, 97% C ₉ H ₈ F ₃ O ₂ , F.W. 206.16, m.p. 32-35°  R:36/37/38, S:26-37		1g	103.00
			5g	415.00
H31659	4-Methoxy-2-(trifluoromethyl)benzyl alcohol, JRD, 97% C ₉ H ₈ F ₃ O ₂ , F.W. 206.16  R:36/37/38, S:26-37		1g	78.00
			5g	312.00
H32993	2-Methoxy-3-(trifluoromethyl)benzylamine, JRD, 97%  C ₉ H ₁₀ F ₃ NO, F.W. 205.18, UN2735  R:34, S:26-36/37/39-45		250mg	54.40
			1g	182.00
H32782	3-Methoxy-5-(trifluoromethyl)benzylamine, JRD, 97%  C ₉ H ₁₀ F ₃ NO, F.W. 205.18, UN2735  R:34, S:26-36/37/39-45		250mg	54.40
			1g	182.00
H32730	4-Methoxy-2-(trifluoromethyl)benzylamine, JRD, 97%  C ₉ H ₁₀ F ₃ NO, F.W. 205.18, UN2735  R:34, S:26-36/37/39-45		1g	127.00
			5g	510.00
H32074	2-Methoxy-3-(trifluoromethyl)benzyl bromide, JRD, 97% C ₉ H ₈ BrF ₃ O, F.W. 269.06, UN3265  R:34, S:26-36/37/39-45		250mg	40.80
			1g	136.00
H31405	2-Methoxy-4-(trifluoromethyl)benzyl bromide, JRD, 97% [886500-59-0], C ₉ H ₈ BrF ₃ O, F.W. 269.06, m.p. 42-44°, UN3261  R:34, S:26-36/37/39-45		1g	100.00
			5g	400.00
H32822	2-Methoxy-6-(trifluoromethyl)benzyl bromide, JRD, 97% C ₉ H ₈ BrF ₃ O, F.W. 269.06, UN3261  R:34, S:26-36/37/39-45		250mg	54.40
			1g	182.00

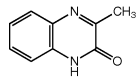
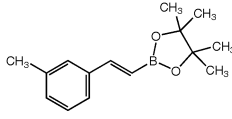
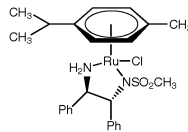
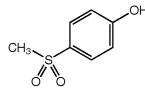
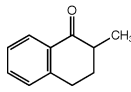
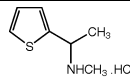
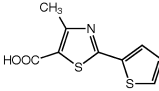
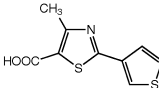
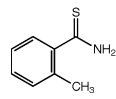
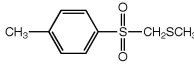
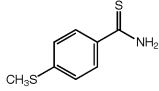
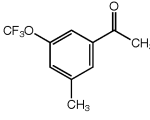
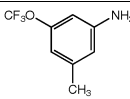
Stock #	Description		Size	\$
H32821	3-Methoxy-5-(trifluoromethyl)benzyl bromide, JRD, 97% C ₉ H ₆ BrF ₃ O, F.W. 269.06, UN3265  R:34, S:26-36/37/39-45		1g	124.00
			5g	497.00
H31871	4-Methoxy-2-(trifluoromethyl)benzyl bromide, JRD, 97% C ₉ H ₆ BrF ₃ O, F.W. 269.06, UN3261  R:34, S:26-36/37/39-45		1g	88.60
			5g	355.00
H31602	2-Methoxy-3-(trifluoromethyl)cinnamic acid, JRD, 97% C ₁₁ H ₈ F ₃ O ₃ , F.W. 246.18  R:36/37/38, S:26-37		250mg	54.40
			1g	182.00
H31803	2-Methoxy-5-(trifluoromethyl)cinnamic acid, JRD, 97% C ₁₁ H ₈ F ₃ O ₃ , F.W. 246.18  R:36/37/38, S:26-37		250mg	33.60
			1g	112.00
H32133	3-Methoxy-5-(trifluoromethyl)cinnamic acid, JRD, 97% C ₁₁ H ₈ F ₃ O ₃ , F.W. 246.18  R:36/37/38, S:26-37		250mg	54.40
			1g	182.00
H32402	4-Methoxy-2-(trifluoromethyl)cinnamic acid, JRD, 97% C ₁₁ H ₈ F ₃ O ₃ , F.W. 246.18  R:36/37/38, S:26-37		250mg	54.40
			1g	182.00
H32777	4-Methoxy-3-(trifluoromethyl)cinnamic acid, JRD, 97% C ₁₁ H ₈ F ₃ O ₃ , F.W. 246.18  R:36/37/38, S:26-37		250mg	54.40
			1g	180.00
H32848	3-Methoxy-5-(trifluoromethyl)phenol, JRD, 97% C ₈ H ₇ F ₃ O ₂ , F.W. 192.14  R:22-36/37/38, S:26-36/37		250mg	54.40
			1g	182.00
H31524	4-Methoxy-3-(trifluoromethyl)phenol, JRD, 97% C ₈ H ₇ F ₃ O ₂ , F.W. 192.14  R:22-36/37/38, S:26-36/37		250mg	54.40
			1g	185.00
H31671	2-Methoxy-3-(trifluoromethyl)phenylacetic acid, JRD, 97% C ₁₀ H ₈ F ₃ O ₃ , F.W. 234.17  R:36/37/38, S:26-37		250mg	67.70
			1g	226.00
H32095	2-Methoxy-6-(trifluoromethyl)phenylacetic acid, JRD, 97% C ₁₀ H ₈ F ₃ O ₃ , F.W. 234.17  R:36/37/38, S:26-37		250mg	97.70
			1g	327.00
H31985	3-Methoxy-5-(trifluoromethyl)phenylacetic acid, JRD, 97% C ₁₀ H ₈ F ₃ O ₃ , F.W. 234.17  R:36/37/38, S:26-37		250mg	95.20
			1g	317.00
H32651	4-Methoxy-2-(trifluoromethyl)phenylacetic acid, JRD, 97% C ₁₀ H ₈ F ₃ O ₃ , F.W. 234.17  R:36/37/38, S:26-37		250mg	67.70
			1g	230.00
H32845	4-Methoxy-3-(trifluoromethyl)phenylacetic acid, JRD, 97% C ₁₀ H ₈ F ₃ O ₃ , F.W. 234.17  R:36/37/38, S:26-37		250mg	67.70
			1g	225.00
H31517	2-Methoxy-3-(trifluoromethyl)phenylacetonitrile, JRD, 97% C ₁₀ H ₇ F ₃ NO, F.W. 215.17, UN3276  R:20/21/22-36/38, S:26-36/37		250mg	54.40
			1g	185.00
H31976	2-Methoxy-6-(trifluoromethyl)phenylacetonitrile, JRD, 97% C ₁₀ H ₇ F ₃ NO, F.W. 215.17, m.p. 48-51°, UN3439  R:20/21/22-36/38, S:26-36/37		250mg	81.70
			1g	273.00

Stock #	Description		Size	\$
H31621	3-Methoxy-5-(trifluoromethyl)phenylacetonitrile, JRD, 97% C ₁₀ H ₈ F ₃ NO, F.W. 215.17, UN3276 ✗ R:20/21/22-36/38, S:26-36/37		250mg 1g	67.70 230.00
H31813	4-Methoxy-2-(trifluoromethyl)phenylacetonitrile, JRD, 97% C ₁₀ H ₈ F ₃ NO, F.W. 215.17, m.p. 30-33°, UN3439 ✗ R:20/21/22-36/38, S:26-36/37		250mg 1g	54.40 182.00
H31730	4-Methoxy-3-(trifluoromethyl)phenylacetonitrile, JRD, 97% C ₁₀ H ₈ F ₃ NO, F.W. 215.17, m.p. 50-53°, UN3439 ✗ R:20/21/22-36/38, S:26-36/37		250mg 1g	54.40 182.00
H32183	3-[2-Methoxy-3-(trifluoromethyl)phenyl]propionic acid, JRD, 97% C ₁₁ H ₁₁ F ₃ O ₃ , F.W. 248.20, m.p. 59-61° ✗ R:36/37/38, S:26-37		250mg 1g	67.70 226.00
H32723	3-[2-Methoxy-5-(trifluoromethyl)phenyl]propionic acid, JRD, 97% C ₁₁ H ₁₁ F ₃ O ₃ , F.W. 248.20 ✗ R:36/37/38, S:26-37		250mg 1g	67.70 230.00
H32338	3-[3-Methoxy-5-(trifluoromethyl)phenyl]propionic acid, JRD, 97% C ₁₁ H ₁₁ F ₃ O ₃ , F.W. 248.20, m.p. 71-73° ✗ R:36/37/38, S:26-37		250mg 1g	67.70 230.00
H32428	3-[4-Methoxy-3-(trifluoromethyl)phenyl]propionic acid, JRD, 97% C ₁₁ H ₁₁ F ₃ O ₃ , F.W. 248.20, m.p. 94-97° ✗ R:36/37/38, S:26-37		250mg 1g	67.70 225.00
H32586	5-Methoxy-2-(trifluoromethyl)pyridine, 97% [216766-13-1], C ₇ H ₆ F ₃ NO, F.W. 177.13, MDL MFCD09864699 ✗ R:36/38, S:26-37		1g 5g	69.00 230.00
H51031	Methyl 3-[4-(aminomethyl)phenyl]propionate hydrochloride C ₁₁ H ₁₅ NO ₂ ·HCl, F.W. 229.71, MDL MFCD11113035 ✗ R:36/37/38, S:26-37		250mg 1g	66.80 241.00
45520	Methyl 4-aminosalicylate, 97% [4-Aminosalicylic acid methyl ester, Methyl 4-amino-2-hydroxybenzoate] [4136-97-4], C ₈ H ₉ NO ₃ , F.W. 167.18, MDL MFCD00088091 ✗ R:22-36/37/38, S:26-36/37		100g 500g	177.00 696.00
H51719	2-Methylbenzamidoxime, 97% [40312-14-9], C ₈ H ₁₀ N ₂ O, F.W. 150.18, m.p. 125-130°, MDL MFCD00052090 ✗ R:36/37/38, S:26-37		1g 5g	45.00 188.00
H51704	4-Methylbenzamidoxime, 97% [19227-13-5], C ₈ H ₁₀ N ₂ O, F.W. 150.18, m.p. 135-138°, RTECS XS4870000, MDL MFCD00019952 ✗ R:36/37/38, S:26-37		1g 5g	48.00 175.00
H51839	5-Methylbenzene-1,3-dithiocarboxamide, 97% C ₈ H ₁₀ N ₂ S ₂ , F.W. 210.31 ✗ R:22-36/37/38, S:26-36/37		250mg 1g	320.00 800.00
H51769	4'-Methylbiphenyl-3-carboxylic acid, 95% [147404-69-1], C ₁₄ H ₁₂ O ₂ , F.W. 212.25 ✗ R:36/37/38, S:26-37		1g 5g	40.50 160.00
H32939	Methyl 4-(Boc-amino)cyclohexylacetate, 95% C ₁₄ H ₂₅ NO ₄ , F.W. 271.35, m.p. 80-82°		100mg 500mg	109.00 365.00
H51905	Methyl 4'-bromobiphenyl-3-carboxylate, 95% C ₁₄ H ₁₁ BrO ₂ , F.W. 291.14		1g 5g	51.60 210.00

Stock #	Description		Size	\$
H51920	Methyl 4'-bromobiphenyl-4-carboxylate, 95% [89901-03-1], C ₁₆ H ₁₁ BrO ₂ , F.W. 291.14, MDL MFCD00496638		1g 5g	54.40 220.00
H31790	Methyl 2-bromo-3-fluoropropionate, 97% C ₅ H ₇ BrFO ₂ , F.W. 184.99, b.p. 50-54°/8mm, UN3265, MDL MFCD09800641		250mg 1g	140.00 390.00
H30222	Methyl 3-bromo-4-hydroxybenzoate, 98% [3-Bromo-4-hydroxybenzoic acid methyl ester] [29415-97-2], C ₈ H ₇ BrO ₃ , F.W. 231.05, m.p. 108-110°		10g 50g	38.80 155.00
H51724	Methyl 3-bromo-5-hydroxybenzoate, 97% [192810-12-1], C ₈ H ₇ BrO ₃ , F.W. 231.05, m.p. 130-135°		250mg 1g	67.50 216.00
H51913	Methyl 3'-bromo-4'-methoxybiphenyl-4-carboxylate, 95% C ₁₅ H ₁₃ BrO ₃ , F.W. 321.17		1g 5g	48.80 195.00
45490	Methyl cellulose [9004-67-5], Powder, Viscosity 15 cPs, MDL MFCD00081763, †		100g 500g 2.5kg	22.30 74.20 269.00
H51910	Methyl 4'-chlorobiphenyl-4-carboxylate, 95% [89901-02-0], C ₁₄ H ₉ ClO ₂ , F.W. 246.69, MDL MFCD04117794		1g 5g	48.80 195.00
H51747	Methyl 5-(4-cyanophenyl)-2-furoate, 95% [834884-75-2], C ₁₃ H ₉ NO ₃ , F.W. 227.22, MDL MFCD06797401		1g 5g	54.40 220.00
H32113	Methyl 4,5-dibromothiophene-2-carboxylate, 97% [62224-24-2], C ₆ H ₄ Br ₂ O ₂ S, F.W. 299.97, m.p. 77-81°		1g 5g	65.60 219.00
H51092	Methyl 2,6-dichloronicotinate, 97% [65515-28-8], C ₇ H ₅ Cl ₂ NO ₂ , F.W. 206.03, MDL MFCD07369794		250mg 1g	58.50 166.00
H30928	Methyl 2-diphenylphosphino-1-naphthoate, 98% [2-Diphenylphosphino-1-naphthoic acid methyl ester] [178176-78-8], C ₂₄ H ₁₉ O ₂ P, F.W. 370.39, m.p. 107-108°		1g 5g 25g	48.50 169.00 657.00
H51781	2-Methyl-4,5-diphenylthiazole, 97% [3755-83-7], C ₁₆ H ₁₃ NS, F.W. 251.35, EINECS 223-161-0, MDL MFCD00186396		1g 5g	57.60 214.00
H30458	Methyl elaidate, 95% [Elaidic acid methyl ester, trans-9-Octadecenoic acid methyl ester] [1937-62-8], C ₁₉ H ₃₅ O ₂ , F.W. 296.50, m.p. 9-10°, d. 0.871, EINECS 217-712-4		1g 10g	44.80 280.00
H51835	3,4-(Methylenedioxy)thiobenzamide, 97% [15884-65-8], C ₈ H ₇ NO ₂ S, F.W. 181.21, MDL MFCD02677739		250mg 1g	39.00 134.00
H31282	5-Methylfuran-2-boronic acid pinacol ester, 97% [338998-93-9], C ₁₁ H ₁₇ BO ₃ , F.W. 208.06, d. 1.023		1g 5g	34.80 116.00
H32829	2-Methylhexanoyl chloride, 97% [41693-47-4], C ₇ H ₁₃ ClO, F.W. 148.63, b.p. 163°, f.p. 55°(131°F), d. 0.957, n _D ²⁰ 1.4290, UN2920		1g 10g	46.70 295.00
H30325	Methyl hydrogen suberate, 98% [mono-Methyl suberate, Suberic acid monomethyl ester] [3946-32-5], CH ₃ O ₂ C(CH ₂) ₆ CO ₂ H, F.W. 223.64, m.p. 17-19°, b.p. 185-186°/18mm, d. 1.04		1g 5g 25g	54.90 150.00 525.00
H51770	Methyl 4'-hydroxybiphenyl-3-carboxylate, 95% [192376-76-4], C ₁₄ H ₁₀ O ₃ , F.W. 228.25, UN3077, MDL MFCD06409246		1g 5g	60.00 240.00

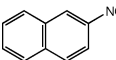
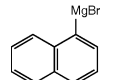
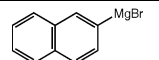
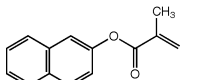
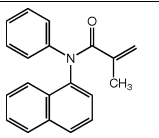
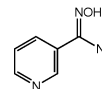
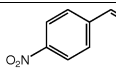
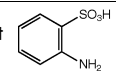
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H32731	Methyl 4-(hydroxymethyl)pyridine-2-carboxylate, 95% [317335-15-2], C ₆ H ₆ NO ₃ , F.W. 167.16, m.p. 74-76° ✗ R:36/37/38, S:26-37		250mg 1g 5g	52.50 145.00 485.00
H51859	1-Methylimidazole-2-carboxylic acid hydrate, 90+% ■ [20485-43-2], C ₅ H ₆ N ₂ O ₃ ·xH ₂ O, F.W. 126.11(anhy), m.p. 116-118°, MDL MFCD01863421 ✗ R:36/37/38, S:26-37		250mg 1g	57.00 182.00
H51866	1-Methylimidazole-4-carboxylic acid ■ [41716-18-1], C ₅ H ₆ N ₂ O ₃ , F.W. 126.11, m.p. 241-243°, MDL MFCD02179560 ✗ R:36/37/38, S:26-37		250mg 1g	60.00 192.00
H51106	1-Methylimidazole-5-carboxylic acid, 95% [41806-40-0], C ₅ H ₆ N ₂ O ₃ , F.W. 126.11, MDL MFCD00955677 ✗ R:36/37/38, S:26-37		250mg 1g	61.90 172.00
H31069	Methyl isocynoacetate, 95% [Isocynoacetic acid methyl ester] [39687-95-1], CNCH ₂ CO ₂ CH ₃ , F.W. 99.09, b.p. 75-76°/10mm, f.p. 84°(183°F), d. 1.09, UN2922, EINECS 254-593-8 ✗ R:20/21/22-34, S:26-36/37/39-45		1g 5g	23.70 67.60
H51924	Methyl 4'-methoxybiphenyl-4-carboxylate, 95% [729-17-9], C ₁₅ H ₁₄ O ₃ , F.W. 242.27		1g 5g	48.80 195.00
H51911	Methyl 4'-methylbiphenyl-4-carboxylate, 95% [49742-56-5], C ₁₅ H ₁₄ O ₂ , F.W. 226.27, EINECS 256-454-7, †		1g 5g	48.80 195.00
H51926	Methyl 4'-(4-morpholinylmethyl)biphenyl-4-carboxylate, 95% C ₁₉ H ₂₁ NO ₃ , F.W. 311.38 ✗ R:36/37/38, S:26-37		1g 5g	58.10 230.00
H31332	2-Methyl-4-nitrobenzoic acid, 97% [1975-51-5], C ₈ H ₇ NO ₄ , F.W. 181.14, m.p. 150-154°, EINECS 217-828-5		1g 5g	61.60 205.00
H30860	3-Methyl-4-nitro-1H-pyrazole, 97% [5334-39-4], C ₅ H ₅ N ₃ O ₂ , F.W. 127.10, m.p. 131-134°, b.p. 325 ✗ R:36/37/38, S:26-37		5g 25g	63.80 212.00
H30430	2-Methylnonanoic acid, 98% [24323-21-5], C ₁₀ H ₂₀ O ₂ , F.W. 172.27, b.p. 265°, UN3265, EINECS 246-162-8, † ✗ R:34, S:26-36/37/39-45		1g 5g	91.50 370.00
H31358	Methyl oleate, 96% [Methyl cis-9-octadecenoate] [112-62-9], CH ₃ (CH ₂) ₇ CH=CH(CH ₂) ₇ CO ₂ CH ₃ , F.W. 296.50, b.p. 168-170°/2mm, d. 0.876, EINECS 203-992-5, RTECS RK0895000		25g 100g	67.50 187.00
H51706	4-(3-Methyl-1,2,4-oxadiazol-5-yl)piperidine hydrochloride, 98% C ₉ H ₁₃ N ₃ O·HCl, F.W. 203.66 ✗ R:36/37/38, S:26-37		250mg 1g	87.30 270.00
H51685	1-(2-Methylphenyl)homopiperazine monohydrochloride, 98% C ₁₂ H ₁₈ N ₂ ·HCl, F.W. 226.75, m.p. 184-188°, Note: Sold in collaboration with Tosoh Organic Chemical Co., Ltd.		500mg	390.00
H51700	1-(3-Methylphenyl)homopiperazine monohydrochloride, 98% [934991-97-6], C ₁₂ H ₁₈ N ₂ ·HCl, F.W. 226.75, m.p. 136-140°, Note: Sold in collaboration with Tosoh Organic Chemical Co., Ltd.		500mg	390.00
H51750	1-(4-Methylphenyl)homopiperazine monohydrochloride, 98% C ₁₂ H ₁₈ N ₂ ·HCl, F.W. 226.75, m.p. 177-181°, Note: Sold in collaboration with Tosoh Organic Chemical Co., Ltd.		500mg	390.00
H51777	5-Methyl-4-phenyl-2-(2-pyridyl)thiazole, 97% C ₁₅ H ₁₂ N ₂ S, F.W. 252.33, m.p. 84-88° ✗ R:36/37/38, S:26-37		1g 5g	63.40 236.00

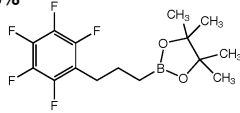
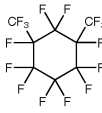
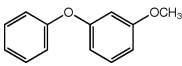
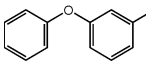
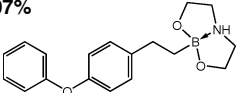
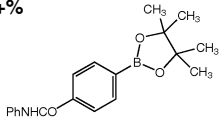
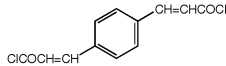
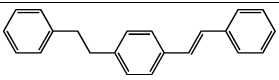
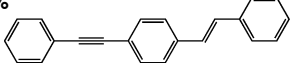
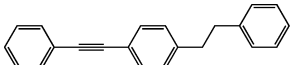
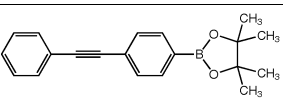
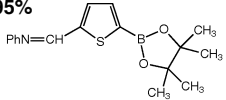
Stock #	Description		Size	\$
H51784	5-Methyl-4-phenyl-2-(3-pyridyl)thiazole, 97% C ₁₅ H ₁₂ N ₂ S, F.W. 252.33, m.p. 77-82° R:36/37/38, S:26-37		1g	63.40
			5g	236.00
H51787	5-Methyl-4-phenyl-2-(4-pyridyl)thiazole, 97% C ₁₅ H ₁₂ N ₂ S, F.W. 252.33, m.p. 82-86° R:36/37/38, S:26-37		1g	63.40
			5g	236.00
H51038	2-Methyl-2-phenylsuccinimide, 99% [1497-17-2], C ₁₁ H ₁₁ NO ₂ , F.W. 189.21, EINECS 216-091-7, MDL MFCD00005496 R:36/37/38, S:26-37		1g	112.00
			5g	453.00
H51827	5-Methyl-4-phenyl-1,2,3-thiadiazole, 97% [64273-28-5], C ₉ H ₈ N ₂ S, F.W. 176.24, MDL MFCD00100201 R:36/37/38, S:26-37		1g	103.00
			5g	383.00
H51728	2-Methyl-4-phenylthiazole, 97% [1826-16-0], C ₁₀ H ₉ NS, F.W. 175.25, m.p. 51-56°		1g	48.00
			5g	175.00
H51659	4-(4-Methyl-1-piperazinyl)benzeneboronic acid pinacol ester [747413-21-4], C ₁₇ H ₂₇ BN ₂ O ₂ , F.W. 302.22		1g	49.90
			5g	198.00
H51786	S-Methyl pyridine-2-carbothioimide hydriodide, 96% ▲ [96898-29-2], C ₇ H ₈ N ₂ S·HI, F.W. 280.13, m.p. 130-135°, MDL MFCD12026515 R:36/37/38, S:26-37		1g	23.30
			5g	86.80
H51776	S-Methyl pyridine-3-carbothioimide hydriodide, 96% ▲ C ₇ H ₈ N ₂ S·HI, F.W. 280.13, m.p. 101-105°, MDL MFCD12026516 R:36/37/38, S:26-37		1g	23.30
			5g	86.80
H51774	S-Methyl pyridine-4-carbothioimide hydriodide, 96% ▲ C ₇ H ₈ N ₂ S·HI, F.W. 280.13, m.p. 182-186°, MDL MFCD12026517 R:36/37/38, S:26-37		1g	23.30
			5g	86.80
H51808	3-Methylpyridine-2-carboxamidoxime, 97% [690632-33-8], C ₇ H ₉ N ₃ O, F.W. 151.168, m.p. 88-92°, MDL MFCD06200877 R:36/37/38, S:26-37		250mg	78.00
			1g	267.00
H51794	3-Methylpyridine-2-thiocarboxamide, 97% C ₇ H ₈ N ₂ S, F.W. 152.21, m.p. 117-122° R:22-36/37/38, S:26-36/37		250mg	78.00
			1g	267.00
H51711	4-Methyl-2-(2-pyridyl)thiazole-5-carboxylic acid, 97% [34418-48-9], C ₁₀ H ₈ N ₂ O ₂ S, F.W. 220.25, m.p. 224-228°, MDL MFCD01316364 R:36/37/38, S:26-37		1g	55.20
			5g	246.00
H51723	4-Methyl-2-(3-pyridyl)thiazole-5-carboxylic acid, 97% [39091-01-5], C ₁₀ H ₈ N ₂ O ₂ S, F.W. 220.25, m.p. 244-248°, MDL MFCD00171753 R:36/37/38, S:26-37		1g	57.50
			5g	251.00
H51753	4-Methyl-2-(4-pyridyl)thiazole-5-carboxylic acid, 97+% [144060-98-0], C ₁₀ H ₈ N ₂ O ₂ S, F.W. 220.25, m.p. 302-307°, MDL MFCD00171757 R:36/37/38, S:26-37		1g	61.30
			5g	262.00
H31236	Methyl (R)-(-)-2-pyrrolidinone-5-carboxylate, 98% [H-D-Pyr-OMe, Methyl D-pyroglytamate] [64700-65-8], C ₆ H ₉ NO ₃ , F.W. 143.14, b.p. 83°/0.25mm, d. 1.226		1g	60.00
			5g	241.00
H51664	3-Methyl-5-(3-pyrrolidinyl)-1,2,4-oxadiazole hydrochloride C ₇ H ₁₁ N ₃ O·HCl, F.W. 189.64, MDL MFCD11870728 R:36/38, S:26-37		250mg	66.00
			1g	165.00
			5g	683.00

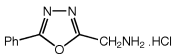
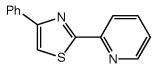
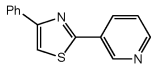
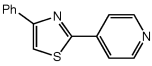
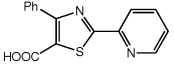
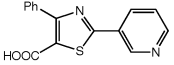
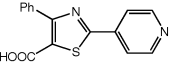
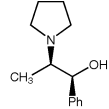
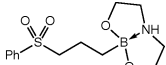
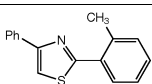
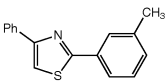
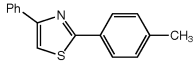
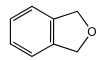
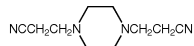
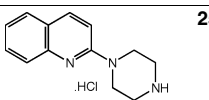
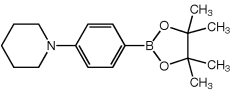
Stock #	Description		Size	\$
H51733	3-Methyl-2(1H)-quinoxalinone, 97% [14003-34-0], C ₉ H ₈ N ₂ O, F.W. 160.18, m.p. 268-273°, RTECS VD3634500		1g 35.70 5g 130.00	
H30073	3-Methyl-β-styrylboronic acid pinacol ester, 97% [4,4,5,5-Tetramethyl-2-(trans-2-m-tolylvinyl)[1,3,2]dioxaborolane] C ₁₅ H ₂₁ BO ₂ , F.W. 244.14, b.p. 114°/0.3mm, n _D ²⁰ 1.5235		1g 54.10 5g 180.00 25g 596.00	
H30016	(R,R)-N-Methylsulfonyl-1,2-diphenylethanediamine(chloro)-(p-cymene)ruthenium(II) [(R,R)-Mesyl-DPEN-(p-cymene)ruthenium chloride] C ₂₅ H ₃₁ ClN ₂ O ₂ RuS, F.W. 560.12 ✗ R:36/37/38, S:26-37		250mg 184.00 1g 550.00	
H32935	4-Methylsulfonylphenol, 97% [14763-60-1], C ₇ H ₈ O ₃ S, F.W. 172.20, m.p. 90-95°, RTECS SM1530000 ✗ R:36/37/38, S:26-37		1g 28.90 10g 180.00	
H31865	2-Methyl-1-tetralone, 98% [1590-08-5], C ₁₁ H ₁₂ O, F.W. 160.22, b.p. 129°/12mm, f.p. >110°(230°F), d. 1.058, n _D ²⁰ 1.5520, EINECS 216-461-8		1g 57.80 10g 385.00	
H51109	N-Methyl-1-(2-thienyl)ethylamine hydrochloride C ₇ H ₉ NS·HCl, F.W. 177.70, MDL MFCD03306017 ✗ R:36/37/38, S:26-37		250mg 64.20 1g 205.00	
H51798	4-Methyl-2-(2-thienyl)thiazole-5-carboxylic acid, 97% C ₉ H ₇ NO ₂ S ₂ , F.W. 225.28 ✗ R:36/37/38, S:26-37		250mg 128.00 1g 384.00	
H51803	4-Methyl-2-(3-thienyl)thiazole-5-carboxylic acid, 97% C ₉ H ₇ NO ₂ S ₂ , F.W. 225.28 ✗ R:36/37/38, S:26-37		250mg 128.00 1g 384.00	
H51790	2-Methyl(thiobenzamide), 97% [53515-19-8], C ₈ H ₈ NS, F.W. 151.23, m.p. 68-73°, MDL MFCD03945350 ✗ R:22, S:36		1g 52.80 5g 197.00	
H32310	Methylthiomethyl acetate, 95% [16437-69-7], CH ₃ CO ₂ CH ₂ SCH ₃ , F.W. 120.17, b.p. 149°, f.p. 44°(111°F), d. 1.086, n _D ²⁰ 1.458, UN3272 ✗ R:10-36/37/38, S:26-37		10g 72.50 50g 241.00	
H31877	Methylthiomethyl p-tolyl sulfone, 99% [59662-65-6], C ₉ H ₁₀ O ₂ S ₂ , F.W. 216.32, m.p. 77-81°		5g 93.30 25g 310.00	
H51848	4-(Methylthio)thiobenzamide, 97% C ₈ H ₈ NS ₂ , F.W. 183.29 ✗ R:36/37/38, S:26-37		1g 57.00 5g 213.00	
H32179	3'-Methyl-5'-(trifluoromethoxy)acetophenone, JRD, 97% C ₁₀ H ₉ F ₃ O ₂ , F.W. 218.17 ✗ R:36/37/38, S:23-26-37		250mg 81.70 1g 273.00	
H30206	3-Methyl-5-(trifluoromethoxy)aniline, JRD, 97% [86256-63-5], C ₈ H ₈ F ₃ NO, F.W. 191.15, b.p. 73°/1mm, n _D ²⁰ 1.4692, UN2810 ✗ R:20/21/22-36/38, S:26-36/37		250mg 83.30 1g 278.00	

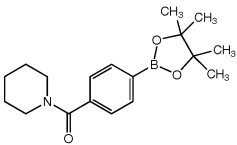
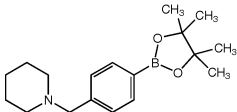
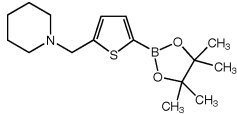
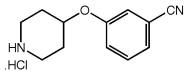
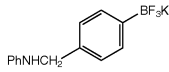
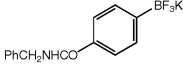
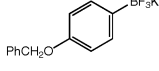
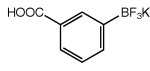
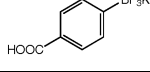
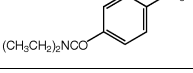
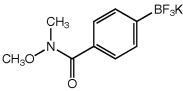
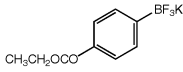
Stock #	Description		Size	\$
H31653	3-Methyl-5-(trifluoromethoxy)benzamide, JRD, 97% C ₉ H ₈ F ₃ NO ₂ , F.W. 219.16 ☒ R:36/37/38, S:26-37		1g 5g	99.20 397.00
H31527	3-Methyl-5-(trifluoromethoxy)benzoic acid, JRD, 97% C ₉ H ₇ F ₃ O ₃ , F.W. 220.15 ☒ R:36/37/38, S:26-37		1g 5g	56.40 226.00
H31921	3-Methyl-5-(trifluoromethoxy)benzonitrile, JRD, 97% C ₉ H ₆ F ₃ NO, F.W. 201.15, UN3276 ☒ R:20/21/22-36/38, S:26-36/37		1g 5g	68.10 273.00
H32689	3-Methyl-5-(trifluoromethoxy)benzoyl chloride, JRD, 97% ▲ C ₉ H ₆ ClF ₃ O ₂ , F.W. 238.59, UN3265 ☒ R:34, S:26-36/37/39-45		1g 5g	68.10 273.00
H32483	3-Methyl-5-(trifluoromethoxy)benzyl alcohol, JRD, 97% C ₉ H ₉ F ₃ O ₂ , F.W. 206.16 ☒ R:36/37/38, S:23-26-37		1g 5g	99.20 397.00
H31840	3-Methyl-5-(trifluoromethoxy)benzylamine, JRD, 97% △ C ₉ H ₁₀ F ₃ NO, F.W. 205.18, UN2735 ☒ R:34, S:26-36/37/39-45		250mg 1g	81.70 273.00
H32444	3-Methyl-5-(trifluoromethoxy)benzyl bromide, JRD, 97% C ₉ H ₈ BrF ₃ O, F.W. 269.06, UN3265 ☒ R:34, S:26-36/37/39-45		1g 5g	115.00 450.00
H31885	3-Methyl-5-(trifluoromethoxy)cinnamic acid, JRD, 97% C ₁₁ H ₉ F ₃ O ₃ , F.W. 246.18 ☒ R:36/37/38, S:26-37		250mg 1g	54.40 182.00
H32888	2'-Methyl-3'-(trifluoromethyl)acetophenone, JRD, 97% C ₁₀ H ₉ F ₃ O, F.W. 202.17 ☒ R:36/38, S:26-37		1g 5g	102.00 411.00
H31819	2'-Methyl-5'-(trifluoromethyl)acetophenone, JRD, 97% C ₁₀ H ₉ F ₃ O, F.W. 202.17 ☒ R:36/38, S:26-37		1g 5g	103.00 411.00
H32226	4'-Methyl-3'-(trifluoromethyl)acetophenone, JRD, 97% C ₁₀ H ₉ F ₃ O, F.W. 202.17 ☒ R:36/38, S:26-37		250mg 1g	67.70 226.00
H31516	2-Methyl-5-(trifluoromethyl)benzamide, 97% [261951-97-7], C ₉ H ₈ F ₃ NO, F.W. 203.17 ☒ R:36/37/38, S:26-37		1g 5g	90.60 365.00
H31552	2-Methyl-3-(trifluoromethyl)benzylamine, JRD, 97% △ C ₉ H ₁₀ F ₃ N, F.W. 189.18, UN2735 ☒ R:34, S:26-36/37/39-45		250mg 1g	54.40 185.00
H32035	Methyl (3-trifluoromethyl)benzylsulfonylacetate, 98% C ₁₁ H ₁₁ F ₃ O ₄ S, F.W. 296.27, m.p. 63-67° ☒ R:36/37/38, S:26-37		250mg 1g 5g	50.90 141.00 471.00
H31937	2-Methyl-5-(trifluoromethyl)cinnamic acid, JRD, 97% C ₁₁ H ₉ F ₃ O ₂ , F.W. 230.18 ☒ R:36/37/38, S:26-37		250mg 1g	40.80 136.00

Stock #	Description		Size	\$
H31502	3-Methyl-5-(trifluoromethyl)cinnamic acid, JRD, 97% C ₁₁ H ₉ F ₃ O ₂ , F.W. 230.18 ✖ R:36/37/38, S:26-37		250mg 1g	54.40 182.00
H31586	2-Methyl-3-(trifluoromethyl)phenol, JRD, 97% C ₈ H ₇ F ₃ O, F.W. 176.14, m.p. 45-48°, UN3261 ✖ R:21/22-34, S:26-36/37/39-45		250mg 1g	54.40 185.00
H31745	2-Methyl-5-(trifluoromethyl)phenol, JRD, 97% C ₈ H ₇ F ₃ O, F.W. 176.14, UN3265 ✖ R:21/22-34, S:26-36/37/39-45		250mg 1g	81.70 273.00
H32051	4-Methyl-3-(trifluoromethyl)phenol, JRD, 97% C ₈ H ₇ F ₃ O, F.W. 176.14, m.p. 45-48° ✖ R:21/22-34, S:26-36/37/39-45		250mg 1g	81.70 273.00
H32357	5-Methyl-2-(trifluoromethyl)phenylacetic acid, JRD, 97% C ₁₀ H ₉ F ₃ O ₂ , F.W. 218.17 ✖ R:36/37/38, S:26-37		250mg 1g	95.20 317.00
H32412	2-Methyl-3-(trifluoromethyl)phenylacetonitrile, JRD, 97% C ₁₀ H ₈ F ₃ N, F.W. 199.17, UN3439 ✖ R:20/21/22-36/38, S:26-36/37		250mg 1g	81.70 275.00
H31940	4-Methyl-3-(trifluoromethyl)phenylacetonitrile, JRD, 97% C ₁₀ H ₈ F ₃ N, F.W. 199.17, UN3276 ✖ R:20/21/22-36/38, S:26-36/37		250mg 1g	67.70 230.00
H32783	5-Methyl-2-(trifluoromethyl)phenylacetonitrile, JRD, 97% C ₁₀ H ₈ F ₃ N, F.W. 199.17, UN3276 ✖ R:20/21/22-36/38, S:26-36/37		250mg 1g	81.70 273.00
H32197	4-Methyl-3-(trifluoromethyl)-DL-phenylalanine, JRD, 97% C ₁₁ H ₁₂ F ₃ NO ₂ , F.W. 247.22		250mg 1g	67.70 226.00
H32002	3-[3-Methyl-5-(trifluoromethyl)phenyl]propionic acid, JRD, 97% C ₁₁ H ₁₁ F ₃ O ₂ , F.W. 232.20, m.p. 53-56° ✖ R:36/37/38, S:26-37		250mg 1g	67.70 220.00
H32500	3-[4-Methyl-3-(trifluoromethyl)phenyl]propionic acid, JRD, 97% C ₁₁ H ₁₁ F ₃ O ₂ , F.W. 232.20 ✖ R:36/37/38, S:26-37		250mg 1g	81.70 273.00
45475	Mini Advanced Coupling Kit Note: Contains 1g each of the following products: 44829 1,1'-Bis(di-tert-butylphosphino)ferrocene palladium dichloride, Pd 16.3%, 45453 Dichlorobis(di-tert-butylphenylphosphine)palladium(II), 10516 Palladium(II) acetate, trimer, Pd 45.9-48.4%, 44618 1,2,3,4,5-Pentaphenyl-1-(di-tert-butylphosphino)ferrocene, CTC-Q-PHOS, 95%, 44730 Palladium anchored homogeneous catalyst, FibreCat™ 1032. This kit contains the most active catalysts and is designed for challenging C-C coupling reactions, amination, alpha ketone arylation, using sterically hindered substrates, aryl chlorides and electron rich substrates. A Catalytic Reaction and Coupling Reference Guide is included. ✖ R:37/38-41, S:26-36/37-38		ea	POR
H51925	4-(4-Morpholinyl)benzeneboronic acid pinacol ester, 95% [568577-88-8], C ₁₆ H ₂₄ BNO ₃ , F.W. 289.18, MDL MFCD04112544 ✖ R:36/37/38, S:26-37		1g 5g	60.00 240.00
H51113	3-(4-Morpholinylcarbonyl)benzeneboronic acid pinacol ester C ₁₇ H ₂₄ BNO ₄ , F.W. 317.19, MDL MFCD09266172 ✖ R:36/37/38, S:26-37		1g 5g 25g	31.70 122.00 492.00
H30191	3-(4-Morpholinyl)propyl-1-boronic acid pinacol ester, 98% [4-[3-(4,4,5,5-Tetramethyl-1,3,2-dioxaborolan-2-yl)propyl]morpholine] C ₁₃ H ₂₅ BNO ₃ , F.W. 255.17 ✖ R:36/37/38, S:26-37		1g 5g	55.30 168.00

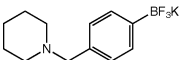
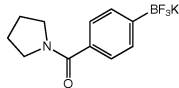
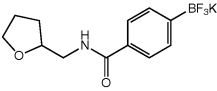
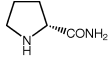
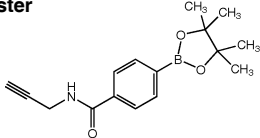
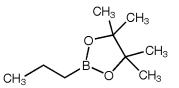
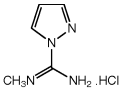
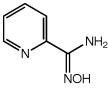
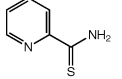
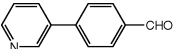
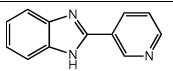
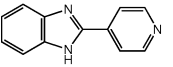
Stock #	Description		Size	\$
H51076	2-Naphthyl isocyanate, 97% ▣ [2243-54-1], C ₁₀ H ₇ NO, F.W. 169.18, m.p. 53-56°, b.p. 83°/0.2mm, UN2811, EINECS 218-815-7, MDL MFCD00014319 ☒ R:20-36/37/38-42, S:22-26-36/37-45		1g	134.00
H51159	1-Naphthylmagnesium bromide, 0.5M in MeTHF ▴ ▣ [703-55-9], C ₁₀ H ₇ BrMg, F.W. 231.37, UN2924, † ☒ R:11-14-19-34, S:8-16-26-36/37/39-43-45		50ml 100ml	112.00 180.00
H51160	2-Naphthylmagnesium bromide, 0.25M in MeTHF ▴ ▣ [21473-01-8], C ₁₀ H ₇ BrMg, F.W. 231.37, UN2924 ☒ R:11-14-19-34, S:8-16-26-36/37/39-43-45		50ml 100ml	115.00 183.00
H51047	2-Naphthyl methacrylate, 95% ▴ [10475-46-4], C ₁₄ H ₁₂ O ₂ , F.W. 212.24, EINECS 233-967-4, MDL MFCD00080581 ☒ R:36/37/38, S:26-28		250mg 1g 5g	49.00 146.00 594.00
H51045	N-(1-Naphthyl)-N-phenylmethacrylamide, 98% ▲ [141029-31-4], C ₂₀ H ₁₇ NO, F.W. 287.36, m.p. 112-118°, MDL MFCD03701557 ☒ R:36/37/38, S:26-37		250mg 1g	70.80 206.00
45566	Neodymium(III) oxide, REACTON®, 99.999% (REO) [1313-97-9], Nd ₂ O ₃ , F.W. 336.48, m.p. 2272°, d. 7.24, Merck 14,6450, EINECS 215-214-1, MDL MFCD00011134, †		25g 100g	177.00 495.00
45498	Nickel sulfide, 99% (metals basis) NiS _{1.4} , UN3077, MDL MFCD00016266 ☒ R:49-43-48/23-68-50/53, S:53-45-60-61		5g 25g	147.00 538.00
H51749	Nicotinamidoxime, 97% [1594-58-7], C ₆ H ₇ N ₃ O, F.W. 137.14, m.p. 130-135°, MDL MFCD00265955 ☒ R:36/37/38, S:26-37		1g 5g	54.00 180.00
45479	Nitric acid, fuming, 98% ■ [7697-37-2], HNO ₃ , F.W. 63.01, d. 1.5, Merck 14,9840, UN2032, EINECS 231-714-2, MDL MFCD00011349, † ☒ R:8-35, S:23-26-36-45		500ml	457.00
H30118	4-Nitrostyrene, 98%, stab. with 3,5-di-tert-butylcatechol [100-13-0], C ₈ H ₇ NO ₂ , F.W. 149.15, m.p. 25-28°, b.p. 93-96°/3mm, f.p. 109°(228°F), d. 1.163, RTECS WL5470000 ☒ R:20/21/22-36/38-68, S:26-36/37		1g 5g	99.00 273.00
H51174	1-Octylmagnesium chloride, 1M in MeTHF ▴ ▣ [38841-98-4], CH ₃ (CH ₂) ₇ CH ₂ MgCl, F.W. 172.98, UN2924, † ☒ R:11-14-19-34, S:8-16-26-36/37/39-43-45		100ml 500ml	52.00 195.00
H31763	Orthanilic acid [88-21-1], C ₈ H ₇ NO ₃ , F.W. 173.19, m.p. >300°, EINECS 201-810-9, RTECS DB4727700, † ☒ R:36/37/38, S:26-37		100g 500g	39.20 136.00
H30179	Oxazole-5-carboxylic acid, 98+% [118994-90-4], C ₄ H ₃ NO ₃ , F.W. 113.07, m.p. ca 245° dec. ☒ R:36/37/38, S:26-37		1g 5g	42.30 141.00
45497	Palladium, 5% on activated carbon powder, Type A103038, sulfided, nominally 50% water wet †		10g 50g 250g	69.00 225.00 825.00
45499	Palladium, 5% on activated carbon powder, Type A405032-5, nominally 50% water wet MDL MFCD03457879, †		5g 25g 100g	41.00 165.00 524.00
45642	Palladium, 5% on silica † ☒ R:48/20, S:36			POR
45515	Palladium, 10% on activated carbon powder, standard, reduced UN1325, †		10g 50g 250g	68.00 269.00 1075.00
45576	Palladium hydroxide, Pd 5% on carbon, nominally 50% water [12135-22-7], EINECS 235-219-2, MDL MFCD00064599, †		2g 10g	39.90 89.50
45512	Palladium(II) nitrate, solution, Pd 4-5% w/w (cont. Pd) [10102-05-3], Pd(NO ₃) ₂ , F.W. 230.43, UN3264, EINECS 233-265-8, MDL MFCD00011169, † ☒ R:34, S:23-26-36/37/39-45		(c)1g (c)5g (c)25g	56.30 225.00 900.00

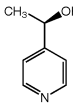
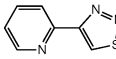
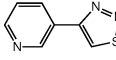
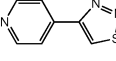
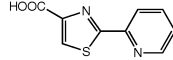
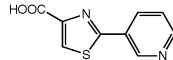
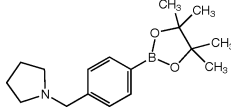
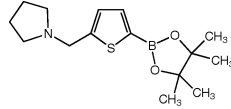
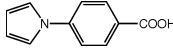
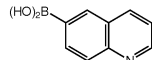
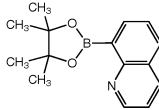
Stock #	Description		Size	\$
H31154	3-Pentafluorophenyl-1-propylboronic acid pinacol ester, 96% [4,4,5,5-Tetramethyl-2-[3-(pentafluorophenyl)propyl]-1,3,2-dioxaborolane] C ₁₃ H ₁₈ BF ₅ O ₂ , F.W. 336.11, b.p. 126-128°/1.5mm, n _D ²⁰ 1.4412  R:36/38, S:26-37		250mg 1g 5g	77.00 214.00 649.00
H51175	1-Pentylmagnesium chloride, 1M in MeTHF   [6393-56-2], CH ₃ (CH ₂) ₄ MgCl, F.W. 130.9, UN2924 R:11-14-19-34, S:8-16-26-36/37/39-43-45		100ml 500ml	54.00 203.00
45477	Perfluoro(1,3-dimethylcyclohexane), tech. 90% [335-27-3], C ₈ F ₁₈ , F.W. 400.06, d. 1.828, MDL MFCD00001469, †		50g 250g	41.70 179.00
H51100	Phenol-d1, 95%(Isotopic)  [1003-66-3], C ₆ H ₅ DO, F.W. 95.12, m.p. 40-42°, b.p. 182°, d. 1.082, UN1671, MDL MFCD01075468  R:23/24/25-34-48/20/21/22-68, S:24/25-26-28-36/37/39-45		1g 5g	88.20 357.00
H32482	3-Phenoxyanisole, 97% [1655-68-1], C ₁₃ H ₁₀ O ₂ , F.W. 200.23, n _D ²⁰ 1.5790, EINECS 216-748-8, RTECS DA5350200, †		5g 25g	46.20 154.00
H32879	3-Phenoxybenzeneboronic acid, 97+% [221006-66-2], C ₁₂ H ₁₁ BO ₃ , F.W. 214.03, m.p. 138-143°  R:36/37/38, S:26-37		250mg 1g	75.60 210.00
H31401	2-(4-Phenoxyphenyl)ethylboronic acid diethanolamine ester, 97% C ₁₈ H ₂₂ BNO ₃ , F.W. 311.19  R:36/37/38, S:26-37		1g 5g 25g	54.10 180.00 595.00
H51668	4-(Phenylaminocarbonyl)benzeneboronic acid pinacol ester, 97+% C ₁₉ H ₂₂ BNO ₃ , F.W. 329.19		1g 5g	32.00 120.00
H51055	1,4-Phenylenediacryloyl chloride, tech.  [35288-49-4], C ₁₂ H ₈ Cl ₂ O ₂ , F.W. 255.10, UN3261, MDL MFCD01109676  R:34, S:26-36/37/39-45		250mg 1g 5g	45.40 132.00 536.00
H31077	1-(trans-2-Phenylethenyl)-4-(2-phenylethyl)benzene, 97% [4-Phenethyl-trans-stilbene, 4-(trans-β-Styryl)bibenzyl] [95166-77-1], C ₂₂ H ₂₀ , F.W. 284.40, m.p. 155-157°		1g	296.00
H30638	1-(trans-2-Phenylethenyl)-4-(phenylethynyl)benzene, 97% [4-(trans-β-Styryl)diphenylacetylene, 4-Ethynyl-trans-stilbene] [21850-30-6], C ₂₂ H ₁₆ , F.W. 280.37		1g	296.00
H30058	1-(2-Phenylethyl)-4-(phenylethynyl)benzene, 97% [4-Phenethyldiphenylacetylene, 4-Phenylethynylbibenzyl] [906650-60-0], C ₂₂ H ₁₈ , F.W. 282.39		1g	296.00
H51699	4-(Phenylethynyl)benzeneboronic acid pinacol ester, 97% C ₂₀ H ₂₁ BO ₂ , F.W. 304.19		1g 5g	65.10 260.00
H51899	2-(Phenyliminomethyl)thiophene-5-boronic acid pinacol ester, 95% C ₁₇ H ₂₀ BNO ₂ S, F.W. 313.22		1g 5g	60.00 240.00
45770	Phenyllithium, typically 1.9M in di-n-butyl ether [591-51-5], C ₈ H ₈ Li, F.W. 84.05, f.p. 20° (68°F), d. 0.85, UN2924, EINECS 209-720-1, BRN 506502, MDL MFCD00014254, †  R:45-46-11-14-34-52/53, S:53-8-16-43-45-61		25ml 100ml 5x100ml	20.00 45.00 200.00
H51157	Phenylmagnesium chloride, 1M in MeTHF   [100-59-4], C ₈ H ₈ MgCl, F.W. 136.86, f.p. -17° (1.4°F), UN2924, †  R:11-14-19-34, S:8-16-26-36/37/39-43-45		100ml 500ml	29.60 74.00

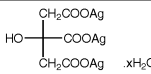
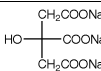
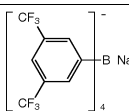
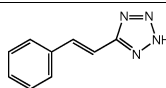
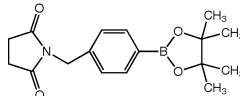
Stock #	Description		Size	\$
H51032	5-Phenyl-1,3,4-oxadiazole-2-methylamine hydrochloride C ₉ H ₉ N ₃ O·HCl, F.W. 211.65, MDL MFCD09028173 ✖ R:36/37/38, S:26-37		1g 5g	57.40 234.00
H51731	4-Phenyl-2-(2-pyridyl)thiazole, 97% [14384-67-9], C ₁₄ H ₁₀ N ₂ S, F.W. 238.31, m.p. 55-59° ✖ R:36/37/38, S:26-37		1g 5g	78.90 288.00
H51748	4-Phenyl-2-(3-pyridyl)thiazole, 97% [70031-86-6], C ₁₄ H ₁₀ N ₂ S, F.W. 238.31, m.p. 52-56° ✖ R:36/37/38, S:26-37		1g 5g	78.90 288.00
H51754	4-Phenyl-2-(4-pyridyl)thiazole, 97% [106950-18-9], C ₁₄ H ₁₀ N ₂ S, F.W. 238.31, m.p. 91-95° ✖ R:36/37/38, S:26-37		1g 5g	82.30 300.00
H51850	4-Phenyl-2-(2-pyridyl)thiazole-5-carboxylic acid, 97% C ₁₅ H ₁₀ N ₂ O ₂ S, F.W. 282.32 ✖ R:36/37/38, S:26-37		250mg 1g	33.00 106.00
H51819	4-Phenyl-2-(3-pyridyl)thiazole-5-carboxylic acid, 97% C ₁₅ H ₁₀ N ₂ O ₂ S, F.W. 282.32 ✖ R:36/37/38, S:26-37		250mg 1g	33.00 106.00
H51826	4-Phenyl-2-(4-pyridyl)thiazole-5-carboxylic acid, 97% C ₁₅ H ₁₀ N ₂ O ₂ S, F.W. 282.32 ✖ R:36/37/38, S:26-37		250mg 1g	33.00 106.00
H31244	(1S,2R)-1-Phenyl-2-(1-pyrrolidinyl)-1-propanol, 98% [(1S,2R)-N-Pyrrolidinylnorephedrine base, (1S,2R)-N,N-Tetramethylenorephedrine] [123620-80-4], C ₁₃ H ₁₉ NO, F.W. 205.30, m.p. 44-46°, b.p. 155°/3mm ✖ R:36/37/38, S:26-37		1g	53.50
H31838	3-(Phenylsulfonyl)propylboronic acid diethanolamine ester C ₁₃ H ₂₀ BNO ₄ S, F.W. 297.18, m.p. 164-166° ✖ R:36/37/38, S:26-37		1g 5g	44.80 150.00
H51828	4-Phenyl-2-(o-tolyl)thiazole, 97% C ₁₆ H ₁₃ NS, F.W. 251.35 ✖ R:36/37/38, S:26-37		1g 5g	70.00 259.00
H51717	4-Phenyl-2-(m-tolyl)thiazole, 97% [2227-70-5], C ₁₆ H ₁₃ NS, F.W. 251.35, m.p. 45-50° ✖ R:36/37/38, S:26-37		1g 5g	78.90 288.00
H51730	4-Phenyl-2-(p-tolyl)thiazole, 97% [2227-61-4], C ₁₆ H ₁₃ NS, F.W. 251.35, m.p. 56-59° ✖ R:36/37/38, S:26-37		1g 5g	82.30 300.00
45592	Phthalan, 96% [1,3-Dihydrobenzo[c]furan, 1,3-Dihydroisobenzofuran] [496-14-0], C ₈ H ₆ O, F.W. 120.15, b.p. 192°, f.p. 63°(145°F), d. 1.094, n _D ²⁰ 1.5460, EINECS 207-815-2, MDL MFCD00005932		5g 25g 100g	30.00 114.00 332.00
H30286	1,4-Piperazinedipropionitrile, 96% [1,4-Bis(2-cyanoethyl)piperazine, 3,3'-Piperazine-1,4-diylbis(propionitrile)] [4159-11-9], C ₁₀ H ₁₆ N ₄ , F.W. 192.27, m.p. 63-66°, UN3439, EINECS 223-992-9 ✖ R:20/21/22-36/37/38, S:26-36/37		1g 5g	75.20 250.00
H51030	2-(1-Piperazinyl)quinoline hydrochloride C ₁₃ H ₁₅ N ₃ ·HCl, F.W. 249.74, MDL MFCD11113034 ✖ R:22-36/37/38, S:26-36/37		250mg 1g	61.80 225.00
H51922	4-(1-Piperidiny)benzeneboronic acid pinacol ester, 95% [852227-96-4], C ₁₇ H ₂₆ BNO ₂ , F.W. 287.21, MDL MFCD07368524		1g 5g	60.00 240.00

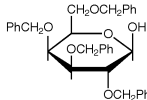
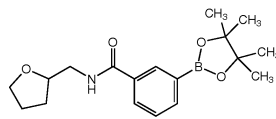
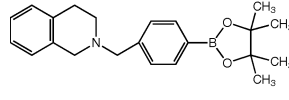
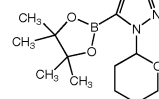
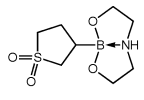
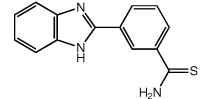
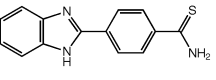
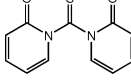
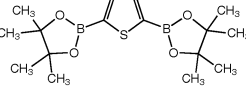
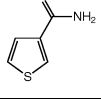
Stock #	Description	Size	\$
H51111	4-(1-Piperidinylcarbonyl)benzeneboronic acid pinacol ester C ₁₈ H ₂₆ BNO ₃ , F.W. 315.21, MDL MFCD09027083	1g	30.00
		5g	115.00
		25g	466.00
			
H51114	4-(1-Piperidinylmethyl)benzeneboronic acid pinacol ester C ₁₈ H ₂₆ BNO ₃ , F.W. 301.23, MDL MFCD08271932	1g	53.40
		5g	205.00
			
H51117	5-(1-Piperidinylmethyl)thiophene-2-boronic acid pinacol ester C ₁₈ H ₂₆ BNO ₃ S, F.W. 307.26, MDL MFCD11113036	1g	63.40
		5g	243.00
			
H51143	3-(4-Piperidinylxy)benzonitrile hydrochloride, 97% [950649-07-7], C ₁₂ H ₁₄ N ₂ O·HCl, F.W. 238.70 ✗ R:22-36/37/38, S:26-36/37	250mg	60.00
		1g	215.00
			
45519	Platinum, 1% on charcoal paste, Type 18MA † ✗ R:43, S:24-37	25g	120.00
		100g	418.00
45778	Platinum, 2% on 2.5mm alumina triblobes MDL MFCD00011179, †	5g	45.00
		25g	180.00
45714	Platinum, 5% on silica †		POR
45594	Polyethyleneimine, branched, M.W. 1,200, 99% [9002-98-6], (-NHCH ₂ CH ₂) _x (-N(CH ₂ CH ₂ NH ₂)CH ₂ CH ₂) _y , MDL MFCD00803910 ✗ R:22, S:36	25g	23.00
		100g	75.00
		500g	302.00
H51149	Potassium 4-(anilinomethyl)phenyltrifluoroborate, 97% C ₁₃ H ₁₂ BF ₃ KN, F.W. 289.15 ✗ R:36/37/38, S:26-37	1g	41.70
		5g	160.00
			
H51887	Potassium 4-(benzylaminocarbonyl)phenyltrifluoroborate, 95% C ₁₄ H ₁₂ BF ₃ KNO, F.W. 317.16 ✗ R:36/37/38, S:26-37	1g	36.60
		5g	145.00
			
H51894	Potassium 4-(benzyloxy)phenyltrifluoroborate, 95% [850623-47-1], C ₁₃ H ₁₁ BF ₃ KO, F.W. 290.13 ✗ R:36/37/38, S:26-37	1g	45.00
		5g	180.00
			
H30530	Potassium tert-butoxide, 1.0M in THF △ ▢ [865-47-4], (CH ₃) ₃ COK, F.W. 112.22, f.p. -19° (-2°F), d. 0.903, UN2924, † ☞ R:11-19-34, S:16-26-36/37/39-43-45	100ml	74.90
		500ml	250.00
		2.5L	824.00
H51885	Potassium 3-carboxyphenyltrifluoroborate, 95% [850313-91-6], C ₇ H ₅ BF ₃ KO ₂ , F.W. 228.02, MDL MFCD07784423 ✗ R:36/37/38, S:26-37	1g	43.00
		5g	170.00
			
H51888	Potassium 4-carboxyphenyltrifluoroborate, 95% [850623-38-0], C ₇ H ₅ BF ₃ KO ₂ , F.W. 228.02, MDL MFCD04115737 ✗ R:36/37/38, S:26-37	1g	45.00
		5g	180.00
			
H51892	Potassium 4-(diethylaminocarbonyl)phenyltrifluoroborate, 95% C ₁₁ H ₁₄ BF ₃ KNO, F.W. 283.14 ✗ R:36/37/38, S:26-37	1g	36.60
		5g	145.00
			
H51896	Potassium 4-(N,O-dimethylhydroxylaminocarbonyl)phenyltrifluoroborate, 95% C ₉ H ₁₀ BF ₃ KNO ₂ , F.W. 271.09 ✗ R:36/37/38, S:26-37	1g	60.00
		5g	240.00
			
H51882	Potassium 4-ethoxycarbonylphenyltrifluoroborate, 95% C ₉ H ₉ BF ₃ KO ₂ , F.W. 256.07 ✗ R:36/37/38, S:26-37	1g	36.60
		5g	145.00
			

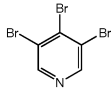
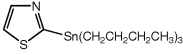
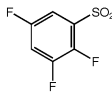
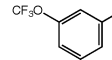
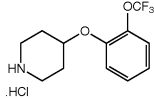
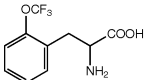
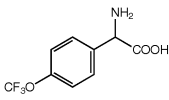
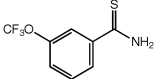
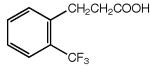
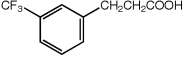
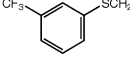
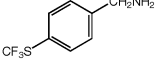
Stock #	Description		Size	\$
H51140	Potassium 5-formylthiophene-2-trifluoroborate Δ [1025113-78-3], $C_5H_4BF_3KOS$, F.W. 218.05, MDL MFCD09800740		1g 5g	48.40 185.00
H51182	Potassium 3-(furfurylaminocarbonyl)phenyltrifluoroborate $C_{12}H_{10}BF_3KNO_2$, F.W. 307.12		1g 5g	41.60 156.00
H51147	Potassium [4-(furfurylaminocarbonyl)phenyl]trifluoroborate $C_{12}H_{10}BF_3KNO_2$, F.W. 307.12, MDL MFCD11977716		1g 5g	41.70 160.00
H51883	Potassium 4-hydroxymethylphenyltrifluoroborate, 95% $C_7H_7BF_3KO$, F.W. 214.04, MDL MFCD08276800		1g 5g	43.00 170.00
H51183	Potassium [3-(2-methoxyethylaminocarbonyl)phenyl-]trifluoroborate $C_{10}H_{12}BF_3KNO_2$, F.W. 285.11, MDL MFCD11977721		1g 5g	38.40 144.00
H51148	Potassium [4-(2-methoxyethylaminocarbonyl)phenyl-]trifluoroborate $C_{10}H_{12}BF_3KNO_2$, F.W. 285.11, MDL MFCD11977717		1g 5g	38.40 147.00
H51151	Potassium 5-methoxypyridine-2-trifluoroborate $C_6H_6BF_3KNO$, F.W. 215.02, MDL MFCD09993103		1g 5g	55.00 206.00
H51889	Potassium 4-(4-methyl-1-piperazinylcarbonyl)phenyl-trifluoroborate, 95% $C_{12}H_{15}BF_3KN_2O$, F.W. 310.17		1g 5g	45.00 180.00
H51884	Potassium 4-(1-methy-4-piperazinyl)phenyltrifluoroborate, 95% $C_{11}H_{15}BF_3KN_2$, F.W. 282.16		1g 5g	72.50 290.00
H51142	Potassium 3-(4-morpholinylcarbonyl)phenyltrifluoroborate $C_{11}H_{12}BF_3KNO_2$, F.W. 297.12, MDL MFCD11977713		1g 5g	53.40 205.00
H51141	Potassium 4-(4-morpholinylcarbonyl)phenyltrifluoroborate $C_{11}H_{12}BF_3KNO_2$, F.W. 297.12, MDL MFCD09800742		1g 5g	53.40 205.00
H51890	Potassium 4-(4-morpholinylmethyl)phenyltrifluoroborate, 95% $C_{11}H_{14}BF_3KNO$, F.W. 283.14		1g 5g	60.00 240.00
H51184	Potassium 3-(phenylaminocarbonyl)phenyltrifluoroborate $C_{13}H_{10}BF_3KNO$, F.W. 303.13, MDL MFCD11977722		1g 5g	29.60 112.00
H51150	Potassium [4-(phenylaminocarbonyl)phenyl]trifluoroborate [912350-01-7], $C_{13}H_{10}BF_3KNO$, F.W. 303.13, m.p. >250°, MDL MFCD11977719		1g 5g	29.60 112.00
H51181	Potassium [3-(1-piperidinylcarbonyl)phenyl]trifluoroborate $C_{12}H_{14}BF_3KNO$, F.W. 295.15		1g 5g	34.50 130.00
H51146	Potassium [4-(1-piperidinylcarbonyl)phenyl]trifluoroborate $C_{12}H_{14}BF_3KNO$, F.W. 295.15, MDL MFCD11977715		1g 5g	40.00 153.00

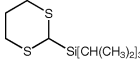
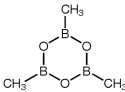
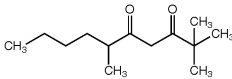
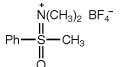
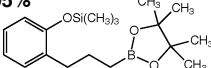
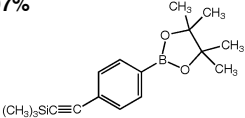
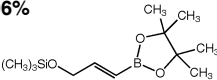
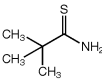
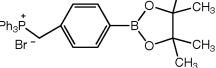
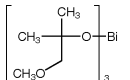
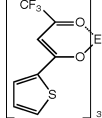
Stock #	Description		Size	\$
H51895	Potassium 4-(1-piperidinylmethyl)phenyltrifluoroborate, 95% C ₁₂ H ₁₆ BF ₃ KN, F.W. 281.17 ✗ R:36/37/38, S:26-37		1g 5g	54.40 220.00
H51893	Potassium 4-(1-pyrrolidinylcarbonyl)phenyltrifluoroborate, 95% C ₁₁ H ₁₂ BF ₃ KNO, F.W. 281.13 ✗ R:36/37/38, S:26-37		1g 5g	36.60 145.00
H51891	Potassium quinoline-8-trifluoroborate, 95% C ₉ H ₆ BF ₃ KN, F.W. 235.06 ✗ R:36/37/38, S:26-37		1g 5g	72.50 290.00
45651	Potassium tantalum isopropoxide, 2.5% w/v in isopropanol   K ₃ Ta[OCH(CH ₃) ₂] ₆ , F.W. 574.57, UN1219 R:11-36/38-67, S:16-26-37		50ml 250ml	58.10 152.00
45548	Potassium tetrabromoplatinate(II), 99.9% (metals basis) [13826-94-3], K ₂ PtBr ₄ , F.W. 592.90, m.p. >300°, EINECS 237-536-1, MDL MFCD00011376, † ✗ R:36/37/38, S:26-37		500mg 2.5g	60.00 250.00
H51886	Potassium 4-(tetrahydrofurfurylaminocarbonyl)phenyltrifluoroborate, 95% C ₁₂ H ₁₄ BF ₃ KNO ₂ , F.W. 311.15 ✗ R:36/37/38, S:26-37		1g 5g	43.00 170.00
H30901	D-(-)-Prolinamide, 98% [62937-45-5], C ₅ H ₁₀ N ₂ O, F.W. 114.10, m.p. 96-102°		1g 5g	50.20 186.00
H51125	4-(Propargylaminocarbonyl)benzeneboronic acid pinacol ester C ₁₆ H ₂₀ BNO ₃ , F.W. 285.15, MDL MFCD11113041		1g 5g	58.40 224.00
H30441	1-Propylboronic acid pinacol ester, 98% [4,4,5,5-Tetramethyl-2-n-propyl-1,3,2-dioxaborolane] [67562-19-0], C ₉ H ₁₈ BO ₂ , F.W. 170.06, b.p. 166-168°, f.p. 47° (117°F), UN1993 R:10		1g 5g 25g	33.30 100.00 361.00
H51162	n-Propylmagnesium chloride, 1M in MeTHF   [2234-82-4], CH ₃ CH ₂ CH ₂ MgCl, F.W. 102.85, UN3399, † R:11-14/15-19-34, S:8-16-26-36/37/39-43-45		100ml 500ml	39.30 148.00
H30607	1H-Pyrazole-1-(N-methylcarboxamidine) hydrochloride, 96% [194852-88-5], C ₅ H ₈ N ₄ ·HCl, F.W. 160.61, m.p. 156-159° ✗ R:36/37/38, S:26-37		5g 25g	63.70 212.00
H51705	Pyridine-2-carboxamidoxime, 97% [1772-01-6], C ₆ H ₇ N ₃ O, F.W. 137.14, m.p. 115-120°, RTECS TJ4310000, MDL MFCD00085159 ✗ R:36/37/38, S:26-37		1g 5g	55.90 193.00
H51710	Pyridine-2-thiocarboxamide, 97+% [5346-38-3], C ₆ H ₆ N ₂ S, F.W. 138.19, m.p. 120-125°, RTECS TJ4305500, MDL MFCD00087576 ✗ R:22-36/37/38, S:26-36/37		5g 25g	25.90 110.00
H51077	4-(3-Pyridyl)benzaldehyde, 98%  [127406-55-7], C ₁₂ H ₉ NO, F.W. 183.21, m.p. 52-57°, MDL MFCD01631910 ✗ R:36/37/38, S:26-37		1g 5g	128.00 519.00
H51785	2-(3-Pyridyl)benzimidazole, 97% [1137-67-3], C ₁₂ H ₉ N ₃ , F.W. 195.22, m.p. 254-260°, MDL MFCD00022665 ✗ R:36/37/38, S:26-37		5g 25g	49.40 188.00
H51778	2-(4-Pyridyl)benzimidazole, 97% [2208-59-5], C ₁₂ H ₉ N ₃ , F.W. 195.22, m.p. 216-221° ✗ R:36/37/38, S:26-37		5g 25g	49.40 188.00

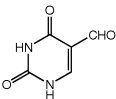
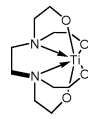
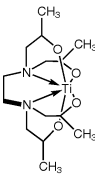
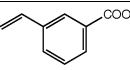
Stock #	Description		Size	\$
H51033	(R)-(+)-1-(4-Pyridyl)ethanol, 99% [27854-88-2], C ₇ H ₉ NO, F.W. 123.15, BRN 4306037, MDL MFCD00077865 ☒ R:36/37/38, S:26-37		250mg 1g	74.00 204.00
H51804	4-(2-Pyridyl)-1,2,3-thiadiazole, 96% C ₇ H ₅ N ₃ S, F.W. 163.20 ☒ R:36/37/38, S:26-37		250mg 1g	58.00 174.00
H51814	4-(3-Pyridyl)-1,2,3-thiadiazole, 96% C ₇ H ₅ N ₃ S, F.W. 163.20 ☒ R:36/37/38, S:26-37		250mg 1g	58.00 174.00
H51821	4-(4-Pyridyl)-1,2,3-thiadiazole, 96% C ₇ H ₅ N ₃ S, F.W. 163.20 ☒ R:36/37/38, S:26-37		250mg 1g	58.00 174.00
H51851	2-(2-Pyridyl)thiazole-4-carboxylic acid, 97% C ₈ H ₆ N ₂ O ₂ S, F.W. 206.22 ☒ R:36/37/38, S:26-37		1g 5g	67.00 249.00
H51838	2-(3-Pyridyl)thiazole-4-carboxylic acid, 97% [39067-29-3], C ₈ H ₆ N ₂ O ₂ S, F.W. 206.22, MDL MFCD00052304 ☒ R:36/37/38, S:26-37		1g 5g	67.00 249.00
45781	Pyrite, naturally occurring mineral, approximately 7.6x7.6cm (3x3in) [1309-36-0], FeS ₂ , F.W. 119.98, d. 5.02, EINECS 215-167-7, MDL MFCD00064690, †		1pc 5pcs	144.00 648.00
H51115	4-(1-Pyrrolidinylmethyl)benzenboronic acid pinacol ester C ₁₇ H ₂₆ BNO ₂ , F.W. 287.20, MDL MFCD08690299		1g 5g	56.70 217.00
H51118	5-(1-Pyrrolidinylmethyl)thiophene-2-boronic acid pinacol ester C ₁₅ H ₁₄ BNO ₂ S, F.W. 293.23, MDL MFCD11113037 ☒ R:36/37/38, S:26-37		1g 5g	65.00 249.00
H51073	4-(1-Pyrrolyl)benzoic acid, 99% ▴ [22106-33-8], C ₁₁ H ₉ NO ₂ , F.W. 187.19, MDL MFCD00082569 ☒ R:36/37/38, S:26-37		1g	103.00
H30380	Quinoline-6-boronic acid hydrochloride, 95% C ₈ H ₆ BNO ₂ ·HCl, F.W. 209.44, m.p. ca 220° dec. ☒ R:36/37/38, S:26-37		250mg 1g	58.40 162.00
H51698	Quinoline-8-boronic acid pinacol ester, 95% [190788-62-6], C ₁₅ H ₁₈ BNO ₂ , F.W. 255.12		1g 5g	65.10 260.00
45538	Rhenium(VII) oxide, 99.995% (metals basis) ▴ [1314-68-7], Re ₂ O ₇ , F.W. 484.40, m.p. 297°, b.p. 360°, d. 6.103, Merck 14,8178, UN3085, EINECS 215-241-9, MDL MFCD00011199, † ☒ R:8-34, S:17-26-36/37/39-45		2g 10g	115.00 350.00
45639	Rhodium, 0.5% on 1mm.(040in) alumina spheres [7440-16-6], EINECS 231-125-0, MDL MFCD00011201, †		10g 50g	95.00 380.00
45674	Ruthenium, 5% on silica † ☒ R:48/20, S:36			POR
45611	Silicon(IV) nitride, amorphous, 98.5+% [12033-89-5], Si ₃ N ₄ , F.W. 140.28, m.p. 1900° subl., d. 3.4, Merck 14,8497, EINECS 234-796-8, MDL MFCD00011230, †		5g 25g 100g	35.20 125.00 399.00
45494	Silver(I) behenate ▴ [Behenic acid silver salt, Sodium docosanoate] [2489-05-6], AgO ₂ C(CH ₂) ₂₀ CH ₃ , F.W. 447.44, MDL MFCD00059001, †		5g 25g 100g	29.90 80.00 290.00
45495	Silver(I) chromate, 99.9% (metals basis) ▴ [7784-01-2], Ag ₂ CrO ₄ , F.W. 331.73, d. 5.63, Merck 14,8510, UN1479, MDL MFCD00003402, † ☒ R:49-8-43-50/53, S:53-45-60-61		5g 25g 100g	22.50 59.50 195.00

Stock #	Description		Size	\$
45496	Silver(I) citrate hydrate ▲ [206986-90-5], C ₆ H ₅ AgO ₇ ·xH ₂ O, F.W. 512.70(anhy), m.p. 170° dec., MDL MFCD00150589, † ☒ R:36/37/38, S:26-37		5g	24.30
			25g	79.90
			100g	275.00
45474	Smopex® -112v Application(s): Recovery of precious metals from catalyst processes, Note: Chopped fibres. Ligand exchanger (thiol functional group). Functional group capacity 2.5mmol/g min		5g	83.00
			25g	282.00
H30611	SnapCure™ 1010 [Titanium polyurethane elastomer catalyst] F.W. 621, d. 1.13, UN1993 ☒ R:10-36/37/38, S:23-26-37		25g	51.70
			100g	86.30
			500g	276.00
H30550	SnapCure™ 1020 ▣ [Zirconium polyurethane elastomer catalyst] F.W. 663, f.p. >80°(176°F), d. 1.07 ☒ R:36, S:23-26		25g	51.70
			100g	86.30
			500g	276.00
H30500	SnapCure™ 1030 ▣ [Zirconium polyurethane elastomer catalyst] F.W. 882, d. 1.14 ☒ R:36/37/38, S:26-37		25g	51.70
			100g	86.30
			500g	276.00
H30790	SnapCure™ 1040 ▣ [Zirconium polyurethane elastomer catalyst] F.W. 1030, f.p. 75°(167°F), d. 1.095		25g	51.70
			100g	86.30
			500g	276.00
H30251	SnapCure™ 3010 ▣ [Titanium polyurethane elastomer catalyst] F.W. 538, f.p. 13°(55°F), d. 1.03, UN1170 ☠☒ R:11-36/38, S:26-37		25g	104.00
			100g	173.00
			500g	553.00
H30922	SnapCure™ 3020 ▣ [Zirconium polyurethane prepolymer catalyst] F.W. 663, f.p. >80°(176°F), d. 1.07 ☒ R:36/37/38, S:26-37		25g	104.00
			100g	173.00
			500g	553.00
H30352	SnapCure™ 3030 ▣ [Zirconium polyurethane prepolymer catalyst] F.W. 867, f.p. 54°(129°F), d. 1.16, UN1993 ☒ R:10-36/37/38, S:26-37		25g	104.00
			100g	173.00
			500g	553.00
45556	Sodium citrate, anhydrous [Citric acid trisodium salt, anhydrous] [68-04-2], C ₆ H ₅ Na ₃ O ₇ , F.W. 258.07, Merck 14,8602, EINECS 200-675-3, RTECS GE8300000, MDL MFCD00012462, †		250g	19.90
			1kg	60.00
45590	Sodium ethoxide, 21% w/w soln. in ethanol, packaged under Argon in resealable ChemSeal™ bottles ▣ [Sodium ethylate] [141-52-6], NaOCH ₂ CH ₃ , F.W. 68.05, f.p. 20°(68°F), d. 0.868, n _D ²⁰ 1.3850, Merck 14,8613, UN3274, EINECS 205-487-5, MDL MFCD00012417, † ☠☒ R:11-14-34, S:8-16-26-43-45		100ml	18.50
			500ml	40.00
45780	Sodium hydroxide, 99.99% (metals basis) △ ▣ [1310-73-2], NaOH, F.W. 40.00, m.p. 316°, d. 2.13, Merck 14,8627, UN1823, EINECS 215-185-5, RTECS WB4900000, MDL MFCD00003548, † ☠☒ R:35, S:26-37/39-45		5g	16.00
			25g	34.80
			100g	72.00
45648	Sodium isopropoxide-isopropanol adduct, 95% △ ▣ NaOC ₃ H ₇ ·C ₃ H ₇ OH, F.W. 148.18, m.p. 150° dec., UN3206 ☠☒ R:11-14-34, S:8-16-26-36/37/39-43-45		5g	25.00
			25g	90.00
		100g	236.00	
H30014	Sodium tetrakis[3,5-bis(trifluoromethyl)phenyl]borate, 97%, may cont.1-5% water [NaBARF24] [79060-88-1], C ₃₂ H ₁₂ BF ₂₄ Na, F.W. 886.20, m.p. 310° dec. ☒ R:36/37/38, S:26-37		1g	105.00
			5g	345.00
			25g	1150.00
45577	Strontium perchlorate trihydrate, 98% ▣ [15650-09-6], Sr(ClO ₄) ₂ ·3H ₂ O, F.W. 340.54 (286.52anhy), UN1508, MDL MFCD09039117, † ☠☒ R:8-36/37/38, S:17-26-37		10g	23.90
			100g	31.90
			500g	95.60
H51107	5-(β-Styryl)-2H-1,2,3,4-tetrazole C ₉ H ₈ N ₄ , F.W. 172.19, MDL MFCD00276928		1g	96.00
			5g	431.00
H51909	4-Succinimidylmethylbenzeneboronic acid pinacol ester, 95% C ₁₇ H ₂₂ BNO ₄ , F.W. 315.17		1g	51.60
			5g	210.00

Stock #	Description	Size	\$	
45596	Sulfuric acid, 72% w/w aq. soln. H ₂ SO ₄ , F.W. 98.08, UN1830, †  R:35, S:26-30-45	1L	85.00	
		4L	156.00	
45542	Sulfuric acid, 15% fuming, 12-17% free SO₃ ■ [Oleum] [8014-95-7], H ₂ SO ₄ + SO ₃ , d. 1.9, Merck 14,8974, UN1831, MDL MFCD00137792, †  R:14-35-37, S:26-30-45	100g	30.10	
		500g 4x500g	76.30 276.00	
45543	Sulfuric acid, 20% fuming, 18-24% free SO₃ ■ [Oleum] [8014-95-7], H ₂ SO ₄ + SO ₃ , d. 1.9, Merck 14,8974, UN1831, MDL MFCD00137792, †  R:14-35-37, S:26-30-45	100g	36.60	
		500g 4x500g	78.50 282.00	
45531	2,3,4,6-Tetra-O-benzyl-D-galactopyranose, 98% [53081-25-7], C ₃₄ H ₃₈ O ₈ , F.W. 540.65, MDL MFCD00132927	 1g 5g	77.00 257.00	
H51670	3-(Tetrahydrofurfurylaminocarbonyl)benzeneboronic acid pinacol ester C ₁₈ H ₂₆ BNO ₄ , F.W. 331.21		1g	29.50
			5g	115.00
H51923	4-(1,2,3,4-Tetrahydro-2-isoquinolinylmethyl)-benzeneboronic acid pinacol ester, 95% C ₂₂ H ₂₈ BNO ₂ , F.W. 349.28  R:36/37/38, S:26-37		1g	54.40
			5g	220.00
H51660	1-(2-Tetrahydropyranyl)-1H-pyrazole-5-boronic acid pinacol ester [903550-26-5], C ₁₄ H ₂₃ BN ₂ O ₃ , F.W. 278.16		1g	50.00
			5g	188.00
H31355	Tetrahydrothiophene-1,1-dioxide-3-boronic acid diethanolamine ester, 97% C ₈ H ₁₆ BNO ₄ S, F.W. 233.10, m.p. 242-244°  R:36/37/38, S:26-37		250mg	93.10
			1g 5g	258.00 878.00
H30637	Thieno[3,2-b]thiophene, 97% [251-41-2], C ₆ H ₄ S ₂ , F.W. 140.23, m.p. 54-58°, b.p. 71°/1mm		1g	154.00
			5g	511.00
H51845	2-(3-Thiocarbamoylphenyl)benzimidazole, 97% C ₁₄ H ₁₁ N ₃ S, F.W. 253.32  R:36/37/38, S:26-37		250mg	88.00
			1g	303.00
H51792	2-(4-Thiocarbamoylphenyl)benzimidazole, 97% C ₁₄ H ₁₁ N ₃ S, F.W. 253.32  R:22-36/37/38, S:26-36/37		250mg	88.40
			1g	303.00
H30087	1,1'-Thiocarbonyldi-2(1H)-pyridone, 95% [102368-13-8], C ₁₁ H ₈ N ₂ O ₂ S, F.W. 232.26, m.p. 163-166°  R:36/37/38, S:26-37		1g	24.10
			5g	82.80
H51725	Thiophene-2,5-diboronic acid bis(pinacol) ester, 97% [175361-81-6], C ₁₈ H ₂₆ B ₂ O ₄ S, F.W. 336.06, m.p. 216-220°		1g	67.50
			5g	270.00
H51830	Thiophene-3-thiocarboxamide, 97% [24044-76-6], C ₅ H ₅ NS ₂ , F.W. 143.22, UN2811, MDL MFCD00173746  R:25-36/37/38, S:26-36/37-45		250mg	88.00
			1g	303.00
45480	Tin(IV) isopropoxide, 99% (metals basis), 10% w/v in isopropanol [1184-61-8], Sn[OCH(CH ₃) ₂] ₄ , F.W. 355.06, UN1993, MDL MFCD00145408   R:11-38-41-67, S:7-16-24/25-26-39		5ml	23.30
			25ml	83.40
			100ml	225.00
			500ml	873.00

Stock #	Description	Size	\$
45603	Titanium(IV) oxide, anatase, 99.7% (metals basis) [1317-70-0], TiO ₂ , F.W. 79.90, Merck 14,9472, EINECS 215-282-2, MDL MFCD00011269, † ☒ R:40, S:36/37	25g 100g 500g	57.00 129.00 441.00
H31732	1H-1,2,4-Triazole-1-acetic acid, 97% [28711-29-7], C ₄ H ₅ N ₃ O ₂ , F.W. 127.10, m.p. 201-205° ☒ R:36/37/38, S:26-37	 250mg 1g	61.10 170.00
H31426	3,4,5-Tribromopyridine, 97% [2457-48-9], C ₅ H ₃ Br ₃ N, F.W. 315.79, m.p. 105-109° ☒ R:21/22-36/37/38, S:26-36/37	 250mg 1g	79.10 220.00
H51090	2-(Tri-n-butylstannyl)thiazole, 96% [121359-48-6], C ₁₅ H ₂₉ NSSn, F.W. 374.17, b.p. 307-309°, d. 1.19, UN2788, MDL MFCD01319057 ☒ R:21-25-36/38-48/23/25-50/53, S:35-36/37/39-45-60-61	 1g 5g	105.00 444.00
H32331	2,3,5-Trifluorobenzenesulfonyl chloride, JRD, 97% ▴ C ₆ H ₂ ClF ₃ O ₂ S, F.W. 230.59, m.p. 47-50°, UN3261 ☒ R:34, S:26-36/37/39-45	 250mg 1g	67.70 230.00
H32174	3-(Trifluoromethoxy)benzenesulfonamide, JRD, 97% C ₇ H ₆ F ₃ NO ₃ S, F.W. 241.19 ☒ R:36/37/38, S:26-37	 1g 5g	68.10 273.00
H51178	4-[2-(Trifluoromethoxy)phenoxy]piperidine hydrochloride C ₁₂ H ₁₄ F ₃ NO ₂ ·HCl, F.W. 297.71, MDL MFCD11870723 ☒ R:36/37/38, S:26-37	 250mg 1g	84.20 268.00
H32667	2-(Trifluoromethoxy)-DL-phenylalanine, JRD, 97% C ₁₀ H ₁₀ F ₃ NO ₃ , F.W. 249.19	 250mg 1g	67.70 230.00
H31245	4-(Trifluoromethoxy)-DL-phenylglycine, 97% [2-Amino-2-(4-trifluoromethoxyphenyl)acetic acid] [261952-24-3], C ₉ H ₉ NO ₃ F ₃ , F.W. 235.16, m.p. ca 292° subl.	 250mg 1g	54.40 180.00
H32377	3-(Trifluoromethoxy)thiobenzamide, 97% C ₈ H ₆ F ₃ NOS, F.W. 222.10, m.p. 76-80° ☒ R:22-36/37/38, S:26-36/37	 250mg 1g	48.80 137.00
H31876	3-[2-(Trifluoromethyl)phenyl]propionic acid, JRD, 97% [94022-99-8], C ₁₀ H ₈ F ₃ O ₂ , F.W. 218.17, † ☒ R:36/37/38, S:26-37	 1g 5g	56.40 226.00
H32088	3-[3-(Trifluoromethyl)phenyl]propionic acid, JRD, 97% C ₁₀ H ₈ F ₃ O ₂ , F.W. 218.17 ☒ R:36/37/38, S:26-37	 1g 5g	51.10 205.00
H51135	[3-(Trifluoromethyl)phenylthio]acetic acid [349-83-7], C ₈ H ₇ F ₃ O ₂ S, F.W. 236.21, MDL MFCD00052114 ☒ R:36/37/38, S:26-37	 1g 5g	80.00 361.00
H31515	4-(Trifluoromethylthio)benzylamine, 95% ▴ [128273-56-3], C ₈ H ₈ F ₃ NS, F.W. 207.21, b.p. 224-225°, n _D ²⁰ 1.5039, UN2735, MDL MFCD00190129 ☒ R:34, S:23-26-36/37/39-45	 1g 5g	122.00 405.00
H51099	1,1,1-Trifluoro-2-phenyl-3-butyne-2-ol, 96% [99727-20-5], C ₁₀ H ₇ F ₃ O, F.W. 200.16, MDL MFCD00792447	 1g 5g	117.00 482.00
H32227	2-(2,3,4-Trifluorophenyl)ethanol, JRD, 90+% C ₈ H ₇ F ₃ O, F.W. 176.14	 250mg 1g	33.60 112.00
H32686	2-(2,4,5-Trifluorophenyl)ethanol, JRD, 97% [883267-70-7], C ₈ H ₇ F ₃ O, F.W. 176.14, MDL MFCD06660356	 250mg 1g	34.70 115.00

Stock #	Description		Size	\$
H32800	2-(3,4,5-Trifluorophenyl)ethanol, JRD, 97% C ₈ H ₇ F ₃ O, F.W. 176.14		250mg 1g	33.60 115.00
H51690	3,3,3-Trifluoropropionaldehyde hemihydrate [936107-87-8], CF ₃ CH ₂ CHO·0.5H ₂ O, F.W. 121.06 (112.05 anhy), MDL MFCD00156074		1g 5g	99.40 435.00
H51071	2-Triisopropylsilyl-1,3-dithiane, 97% Δ [145251-89-4], C ₁₃ H ₂₈ S ₂ Si, F.W. 276.58, m.p. 46-49°, MDL MFCD01863487  R:36/37/38, S:26-37		1g 5g	138.00 568.00
H31111	2,4,6-Trimethylboroxin, 99% [823-96-1], C ₃ H ₆ B ₃ O ₃ , F.W. 125.54, m.p. -38°, b.p. 78-80°, d. 0.90, UN2924   R:11-34, S:26-36/37/39-45		5g 25g	88.50 350.00
H32926	2,2,6-Trimethyl-3,5-decanedione, 95% C ₁₃ H ₂₄ O ₂ , F.W. 212.18, f.p. >61° (142°F)		250mg 1g	83.10 230.00
H32401	N,N,S-Trimethyl-S-phenylsulfoximinium tetrafluoroborate, tech. 90% [21077-81-6], C ₆ H ₄ NOS BF ₄ ⁻ , F.W. 271.08, m.p. 118-120°, MDL MFCD08741486  R:36/37/38, S:26-37		250mg 1g	120.00 337.00
H30323	3-(2-Trimethylsiloxyphenyl)-1-propylboronic acid pinacol ester, 95% [4,4,5,5-Tetramethyl-2-[3-(2-trimethylsiloxyphenyl)propyl]-1,3,2-dioxaborolane] C ₁₈ H ₃₁ BO ₃ Si, F.W. 334.34, n _D ²⁰ 1.4799		1g 5g 25g	60.80 200.00 662.00
H51697	4-[(Trimethylsilyl)ethynyl]benzeneboronic acid pinacol ester, 97% C ₁₇ H ₂₅ BO ₂ Si, F.W. 300.28		1g 5g	78.80 310.00
H51689	trans-3-Trimethylsiloxy-1-propenylboronic acid pinacol ester, 96% C ₁₂ H ₂₃ BO ₃ Si, F.W. 256.22		1g 5g 25g	70.20 276.00 1150.00
H32092	2,2,2-Trimethylthioacetamide, 97% [630-22-8], C ₆ H ₁₁ NS, F.W. 117.21, m.p. 113-117°  R:22, S:36		1g 10g	47.80 295.00
H51917	4-(Triphenylphosphoniomethyl)benzeneboronic acid pinacol ester bromide, 95% C ₃₁ H ₃₃ BrO ₂ P, F.W. 559.29		1g 5g	48.80 195.00
H51864	Tris(2-carboxyethyl)phosphine hydrochloride, 0.5M soln in water [51805-45-9], (HO ₂ CCH ₂ CH ₂) ₃ P·HCl, F.W. 286.65, MDL MFCD00145469  R:36/38, S:26-37		5ml 25ml 100ml	32.00 135.00 440.00
H31853	Tris(2-dimethylaminoethyl)amine, 99+% [33527-91-2], [(CH ₃) ₂ NCH ₂ CH ₂] ₃ N, F.W. 230.40, b.p. 70-71°/0.5mm, n _D ²⁰ 1.4540, UN2735  R:34, S:26-36/37/39-45		5g 25g	103.00 343.00
45488	Tris(1-methoxy-2-methyl-2-propoxy)bismuth, 99.99% (metals basis) C ₁₅ H ₃₃ O ₆ Bi, F.W. 518.40, m.p. 70°  R:36/37/38, S:26-37		1g 5g 25g	80.60 323.00 1054.00
45772	Tris(4,4,4-trifluoro-1-(2-thienyl)-1,3-butanediono)europium(III) [Europium(III) thenoyltrifluoroacetate] [14054-87-6], C ₂₄ H ₁₂ EuF ₉ O ₆ S ₃ , F.W. 815.51, m.p. 134-140°, EINECS 237-892-8, MDL MFCD00041948, t  R:36/37/38, S:26-37		1g	220.00
45513	Tungsten(V) ethoxide, 94% $\square$ [26143-11-3], W(OCH ₂ CH ₃) ₅ , F.W. 409.15, m.p. <20°, b.p. 110-115°/1mm, f.p. >110° (230°F), MDL MFCD00078041  R:36/38, S:26-37		2g 10g	41.90 172.00

Stock #	Description	Size	\$
45659	Tungsten(V) ethoxide 1,2-dimethoxyethane adduct, 99%   W(OC ₂ H ₅) ₅ C ₄ H ₁₀ O ₂ , F.W. 499.26, m.p. <0°, b.p. dec., UN1993  R:60-61-11-19-20-36/38, S:53-45	5g 25g	48.80 195.00
H51098	Uracil-5-carboxaldehyde, 97%  [1195-08-0], C ₄ H ₃ N ₂ O ₃ , F.W. 140.10, MDL MFCD00192185	 250mg 1g	44.40 148.00
H31066	VERTEC™ LAC1000 [[2,2',2'',2'''-(Ethanediylidinitrilo)tetrakis(ethoxy)-N,N',O,O',O'',O''']titanium(IV), N,N,N',N'-Tetrakis(2-ethoxy)ethylenediamine titanium(IV)] [17476-13-0], C ₁₀ H ₂₀ N ₂ O ₄ Ti, F.W. 280.15, MDL MFCD11113032  R:36/37/38, S:26-37	 5g 25g 100g	44.80 180.00 563.00
H31123	VERTEC™ LAC2000 [[2,2',2'',2'''-(Ethanediylidinitrilo)tetrakis(2-propoxy)-N,N',O,O',O'',O''']titanium(IV), N,N,N',N'-Tetrakis(2-propoxy)ethylenediamine titanium(IV)] [96471-81-7], C ₁₄ H ₂₈ N ₂ O ₄ Ti, F.W. 336.26, MDL MFCD11113033  R:36/37/38, S:26-37	 5g 25g 100g	44.80 180.00 563.00
H31723	2-Vinylbenzoic acid, 96% [27326-43-8], C ₉ H ₈ O ₂ , F.W. 148.16, m.p. 94°  R:36/37/38, S:26-37	 1g 5g	58.10 179.00
H32556	3-Vinylbenzoic acid, 96% [28447-20-3], C ₉ H ₈ O ₂ , F.W. 148.16, m.p. 95°  R:36/37/38, S:26-37	 1g 5g	41.50 128.00
H51043	1-Vinylnaphthalene, 95%, stab. with 4-tert-butylcatechol [826-74-4], C ₁₂ H ₁₀ , F.W. 154.21, EINECS 212-560-5, MDL MFCD00075766  R:36/37/38, S:26-37	 1g 5g	53.90 203.00
45683	Zinc oxide, NanoArc® ZN-2210, 40% in mineral spirits, colloidal dispersion with dispersant [1314-13-2], ZnO, F.W. 81.37, Application(s): Zinc oxide dispersed in mineral spirits suitable for incorporation in a wide variety of non-polar organic based coatings and formulations where high transparency is important, UN1993, MDL MFCD00011300, Note: Sold in collaboration with Nanophase Technologies Corporation  R:10-50/53, S:60-61	100g 500g	26.00 115.00
45632	Zinc oxide, NanoArc® ZN-2225, 40% in 1,2-propanediol monomethyl ether acetate, colloidal dispersion with dispersant [1314-13-2], ZnO, F.W. 81.37, Application(s): Zinc oxide dispersed in Dowanol PMA suitable for incorporation in a wide variety of polar organic based coatings and formulations where high transparency is important, UN1993, MDL MFCD00011300, Note: Sold in collaboration with Nanophase Technologies Corporation  R:10-50/53, S:60-61	100g 500g	24.00 105.00
45646	Zinc oxide, NanoTek® ZN-2525, 40% in 1,2-propanediol monomethyl ether acetate, colloidal dispersion with dispersant [1314-13-2], ZnO, F.W. 81.37, Application(s): Zinc oxide dispersed in Dowanol PMA suitable for incorporation in a wide variety of polar organic based coatings and formulations, UN1993, MDL MFCD00011300, Note: Sold in collaboration with Nanophase Technologies Corporation  R:10-50/53, S:60-61	100g 500g	22.00 95.00
45588	Zinc oxide, NanoTek® ZN-2551, 50% in H₂O, colloidal dispersion with dispersant [1314-13-2], ZnO, F.W. 81.37, Application(s): Water borne zinc oxide dispersion suitable for incorporation in water compatible coatings and formulations, UN3082, MDL MFCD00011300, Note: Sold in collaboration with Nanophase Technologies Corporation  R:50/53, S:60-61	100g 500g	21.00 80.00
45748	Zinc oxide, NanoTek® ZN-2510, 50% in mineral spirits, colloidal dispersion with dispersant [1314-13-2], ZnO, F.W. 81.37, Application(s): Zinc oxide dispersed in mineral spirits suitable for incorporation in a wide variety of non-polar organic based coatings and formulations, UN1993, MDL MFCD00011300, Note: Sold in collaboration with Nanophase Technologies Corporation  R:10-50/53, S:60-61	100g 500g	20.00 90.00
45801	Zinc oxide, NanoTek® ZN-2610, 50% in mineral spirits, colloidal dispersion with dispersant [1314-13-2], ZnO, F.W. 81.37, Application(s): Zinc oxide dispersed in mineral spirits suitable for incorporation in a wide variety of non-polar organic based coatings and formulations, UN1993, MDL MFCD00011300, Note: Sold in collaboration with Nanophase Technologies Corporation  R:10-50/53, S:60-61	100g 500g	21.00 70.00

Stock #	Description	Size	\$
45486	Zinc oxide substrate, 10x10x0.5mm, polished one side, 0001 orientation ZnO, EINECS 215-171-9, MDL MFCD00011109, †  R:50/53, S:60-61	1each	550.00
		5each	1926.00
45487	Zinc oxide substrate, 10x10x0.5mm, polished two sides, 0001 orientation ZnO, EINECS 215-171-9, MDL MFCD00011109, †  R:50/53, S:60-61	1each	925.00
		5each	3144.00
45562	Zirconium(IV) oxide, yttria stabilized, 99.95% (metals basis excluding Hf) [1314-23-4], ZrO ₂ + 8% Y ₂ O ₃ , m.p. ca 2700°, b.p. ca 5000°, Merck 14,10180, EINECS 215-227-2, RTECS ZH8800000, MDL MFCD00145546, †	100g	200.00
		500g	800.00
45600	Zirconium oxide, catalyst support, sulfated m.p. 3500°, d. 5.5, †  R:36/38, S:26-37	100g	83.00
		500g	265.00
		2.5kg	925.00
45597	Zirconium oxide, catalyst support, tungstated †  R:36/37/38, S:26-37	100g	83.00
		500g	265.00
		2.5kg	925.00



Section 2 Pure Elements & Alloys

Alphabetical listing of Pure Elements	79
Alphabetical listing of Alloys	82

This section lists new pure elements in various forms and purities and new alloys in several compositions and forms. Our broad range of products is designed to meet a wide variety of high technology applications requiring pure elements. Different purities are provided to correspond to the demands of many different uses. Representative physical properties and additional facts pertaining to each element are summarized at the top of the page. For fabricated metals (foils, wires, rods, etc.) tolerances are $\pm 10\%$ of the listed value.

All products designated as PuraTronic®, REacton®, or Premion® grade are automatically supplied with a certificate of analysis. For PuraTronic® and Premion® products, the percent purity is based upon total metallic impurities. REacton® products' purities are based on total rare earth oxide (REO) impurities.

If you cannot find the purity, form or particle size you need, simply call our Speciality and Bulk Sales Department.

Glossary of terms describing forms

The various forms appear in the order below. Special items specific to particular elements are noted before powders. For example, in the Gold (Au) listing, Bright Brushing Gold is listed before the powders.

1. Powder - solid material with a very small particle size
2. Flake - powder with a flat, irregular shape
3. Granules - uniform, amorphous pieces of material
4. Spheres - uniform sized, spherical pieces of material
5. Shot - spherical to semi-spherical pieces of material of varying sizes
6. Splatter - pieces formed by dropping molten metal onto a cooling surface
7. Needles - uniform, elongated pieces of material
8. Turnings - small concentric shavings machined from a larger form
9. Sponge - pieces with a high surface area resulting from complex surface morphology
10. Mossy - pieces formed by dropping molten metal into water
11. Lump - a solid piece of amorphous material, larger than a granule
12. Pieces - solid pieces of material, larger than a granule
13. Cubes - uniform sized, cubic shaped pieces of material
14. Sticks - rectangular pieces of a material
15. Bar - a rectangular or cylindrical piece of material
16. Ingot - a cast, usually rectangular piece of material
17. Pellets - somewhat regular shaped pieces of material
18. Slugs - short cylindrical pieces of material of varying lengths and diameters
19. Yarn - a parallel collection of a definite number of fiber strands, usually three to several hundred
20. Felt - compressed, porous, nonwoven fabric
21. Fiber - a pure monofilament form of solid material having an extremely high length to diameter ratio
22. Microleaf - an extremely thin layer of pure material on a temporary support ≤ 1 micron thick
23. Ultrathin foil - an extremely thin sheet of pure material, supported or unsupported ≤ 1 micron thick
24. Thin foil - a very thin sheet of unsupported pure material 1.1 - 24.0 micron thick
25. Foil - a thin sheet of pure material, 0.025mm - 2mm
26. Ribbon - a thin width of foil, offered in rolls of varying length
27. Disc - a cylindrical piece of material with a diameter much larger than the thickness
28. Sputtering target - a disc of high purity material used as an atomic sputtering source for ion bombardment
29. Plate - a sheet of fabricated pure material > 2 mm thick
30. Gauze - a wire cloth material consisting of wires of a pure material woven into a grid having consistent openings
31. Wire - a uniform strand of a material having a diameter ≤ 2.0 mm
32. Rod - a uniform strand of a material having a diameter ≥ 2.0 mm
33. Tubing - a uniform strand of a material having a hollowed core

Pure Elements

Stock #	Description	Size	\$
45546	Aluminum powder, spherical, APS approximately 0.07 micron [7429-90-5], UN1396, MDL MFCD00134029, †  R:11-15, S:7/8-33-43	25g 100g	108.00 333.00
45769	Arsenic polycrystalline lump, 2-8mm (0.08-0.3in), Puratronic®, 99.99999+% (metals basis) [7440-38-2], UN1558, EINECS 213-148-6, RTECS CG0525000, MDL MFCD00085309, †   R:23/25-50/53, S:20/21-28-45-60-61	1g 10g 25g 100g 1kg	65.50 91.40 158.00 393.00 2112.00
45527	Carbon black, acetylene, 100% compressed, 99.5% (metals basis) [1333-86-4], S.A. 60-80m ² /g, Bulk density 8.43-12.9lbs/ft ³ , UN1325, EINECS 215-609-9, RTECS FF5800000, MDL MFCD00133992, †   R:11-40, S:36/37	250g 2x500g	38.20 91.70
45478	Carbon, activated, -20+40 mesh [7440-44-0], EINECS 231-153-3, MDL MFCD00133992, †	100g 500g 2kg	14.30 25.50 78.40
45618	Cerium sputtering target, 50.8mm dia x 1.59mm thick [7440-45-1], Ce, UN1333, EINECS 231-154-9, MDL MFCD00010924, †	each	680.00
45734	Cerium sputtering target, 50.8mm dia x 3.18mm thick [7440-45-1], Ce, UN1333, EINECS 231-154-9, MDL MFCD00010924, †  R:11-15, S:7/8/43	each	720.00
46058	Cerium sputtering target, 76.2mm dia x 1.59mm thick [7440-45-1], Ce, UN1333, EINECS 231-154-9, MDL MFCD00010924, †	each	985.00
45926	Cerium sputtering target, 76.2mm dia x 3.18mm thick [7440-45-1], Ce, UN1333, EINECS 231-154-9, MDL MFCD00010924, †	each	1050.00
45506	Cobalt nanopowder, APS 5-15nm, 99.9% (metals basis) [7440-48-4], Co(core)/Co ₃ O ₄ (shell), Co ₃ O ₄ layer 0.5-1.5nm, UN3089, MDL MFCD00010935, †   R:11-42/43-53, S:22-24-37-61	5g 25g	258.00 860.00
45544	Copper powder, spherical, -170+270 mesh, 99.9% (metals basis) [7440-50-8], UN3089, MDL MFCD00010965, †   R:11-36/37/38, S:26-37	100g 500g 2kg	32.30 108.00 349.00
45504	Copper nanopowder, APS 20-40nm, 99.9% (metals basis) [7440-50-8], Cu(core)/CuO(shell), CuO layer 1-2nm, UN3089, MDL MFCD00010965, †   R:11-36/37, S:26	5g 25g	129.00 430.00
45619	Dysprosium sputtering target, 50.8mm dia x 1.59mm thick [7429-91-6], Dy, EINECS 231-073-9, MDL MFCD00010982, †	each	710.00
45735	Dysprosium sputtering target, 50.8mm dia x 3.18mm thick [7429-91-6], Dy, EINECS 231-073-9, MDL MFCD00010982, †	each	780.00
46059	Dysprosium sputtering target, 76.2mm dia x 1.59mm thick [7429-91-6], Dy, EINECS 231-073-9, MDL MFCD00010982, †	each	1100.00
45927	Dysprosium sputtering target, 76.2mm dia x 3.18mm thick [7429-91-6], Dy, EINECS 231-073-9, MDL MFCD00010982, †	each	1225.00
45623	Europium sputtering target, 50.8mm dia x 1.59mm thick [7440-53-1], Eu, UN3178, EINECS 231-161-7, MDL MFCD00010992, †		POR
45739	Europium sputtering target, 50.8mm dia x 3.18mm thick [7440-53-1], Eu, UN3178, EINECS 231-161-7, MDL MFCD00010992, †		POR
46063	Europium sputtering target, 76.2mm dia x 1.59mm thick [7440-53-1], Eu, UN3178, EINECS 231-161-7, MDL MFCD00010992, †		POR
45931	Europium sputtering target, 76.2mm dia x 3.18mm thick [7440-53-1], Eu, UN3178, EINECS 231-161-7, MDL MFCD00010992, †		POR
45529	Gadolinium ingot, 99.9% (REM) [7440-54-2], ≈40-80g/piece, irreg. shape, packaged under argon, EINECS 231-160-1, MDL MFCD00011022, †	1pc	237.00
45621	Gadolinium sputtering target, 50.8mm dia x 1.59mm thick [7440-54-2], Gd, EINECS 231-160-1, MDL MFCD00011022, †	each	655.00
45737	Gadolinium sputtering target, 50.8mm dia x 3.18mm thick [7440-54-2], Gd, EINECS 231-160-1, MDL MFCD00011022, †	each	700.00
46061	Gadolinium sputtering target, 76.2mm dia x 1.59mm thick [7440-54-2], Gd, EINECS 231-160-1, MDL MFCD00011022, †	each	990.00

Stock #	Description	Size	\$
45929	Gadolinium sputtering target, 76.2mm dia x 3.18mm thick [7440-54-2], Gd, EINECS 231-160-1, MDL MFCD00011022, †	each	1070.00
45567	Gold wire, 0.25mm (0.01in) dia, annealed, Premion®, 99.999% (metals basis) [7440-57-5], ≈0.95g/m, MDL MFCD00003436, †	25cm 100cm 500cm	58.00 175.00 700.00
45507	Iron nanopowder, APS 10-30nm, 99.9% (metals basis) △ [7439-89-6], Fe(core)/FeO(shell), FeO layer 0.5-1.5nm, UN3089, EINECS 231-096-4	5g 25g	226.00 753.00
	  R:11-36/37, S:16-26		
45624	Lanthanum sputtering target, 50.8mm dia x 1.59mm thick [7439-91-0], La, UN3178, EINECS 231-099-0, MDL MFCD00011066, †	each	660.00
45740	Lanthanum sputtering target, 50.8mm dia x 3.18mm thick [7439-91-0], La, UN3178, EINECS 231-099-0, MDL MFCD00011066, †	each	685.00
46064	Lanthanum sputtering target, 76.2mm dia x 1.59mm thick [7439-91-0], La, UN3178, EINECS 231-099-0, MDL MFCD00011066, †	each	930.00
45932	Lanthanum sputtering target, 76.2mm dia x 3.18mm thick [7439-91-0], La, UN3178, EINECS 231-099-0, MDL MFCD00011066, †	each	970.00
45602	Lead powder, -100 mesh, 99.95% (metals basis) [7439-92-1], Pb, atomized powder, UN3077, EINECS 231-100-4, MDL MFCD00134050, †	250g 1kg	28.90 67.00
	  R:61-62-20/22-33-50/53, S:53-45-60-61		
45521	Lead ribbon, 76.2mm (3.0in) wide, 0.15mm (0.006in) thick, low silver (<0.0001%), [7439-92-1], Assay grade foil, ≈132g/m, EINECS 231-100-4, MDL MFCD00134050, †	5lb	85.00
	  R:61-62-33-50/53, S:53-45-60-61		
45508	Manganese nanopowder, APS 30-50nm, 99.9% (metals basis) [7439-96-5], Mn(core)/Mn ₂ O ₄ (shell), Mn ₂ O ₄ layer 0.5-1.5nm, UN3089, EINECS 231-105-1, MDL MFCD00011111	5g 25g	210.00 699.00
	 R:11, S:33		
45505	Nickel nanopowder, APS 5-20nm, 99.9% (metals basis) [7440-02-0], Ni(core)/NiO(shell), NiO layer 0.5-1.5nm, UN3089, EINECS 231-111-4	5g 25g	258.00 860.00
	  R:11-40-43-48/23-52/53, S:36/37/39-45-61		
45524	Rhenium slug, 12.7mm (0.5in) dia, 99.99% (metals basis) [7440-15-5], ≈6.35mm long, 7.5 +/- 2 grams, MDL MFCD00011195, †	1each 5each	376.00 1371.00
45484	Ruthenium cubes, 6mm (0.24in) square, 99.9% (metals basis) [7440-18-8], ≈2-3g/cube, MDL MFCD00011207, †	1pc	495.00
45609	Silicon powder, crystalline, -100+200 mesh, 99.99% (metals basis) [7440-21-3], UN3089, EINECS 231-130-8, MDL MFCD00085311, †	10g 50g 250g	35.00 80.00 210.00
	  R:11-36/37/38, S:26-36/37/39		
45599	Silicon powder, crystalline, APS ≤50nm, 99+%, Plasma Synthesized [7440-21-3], Packed under Argon, UN3089, EINECS 231-130-8, MDL MFCD00085311, †	2g 10g 50g	36.00 145.00 650.00
	  R:11-36/37/38, S:26-37		
45661	Silver Conductive Ink S-020 (formerly Mattheylec P2006), d. 2.5	5g 25g 100g	15.00 60.00 200.00
45509	Silver nanopowder, APS 20-40nm, 99.9% (metals basis) [7440-22-4], Ag, No Oxide layer, MDL MFCD00003397, †	5g 25g	135.00 450.00
45622	Terbium sputtering target, 50.8mm dia x 1.59mm thick [7440-27-9], Tb, EINECS 231-137-6, MDL MFCD00011256, †	each	815.00
45738	Terbium sputtering target, 50.8mm dia x 3.18mm thick [7440-27-9], Tb, EINECS 231-137-6, MDL MFCD00011256, †	each	920.00
46062	Terbium sputtering target, 76mm dia x 1.59mm thick [7440-27-9], Tb, EINECS 231-137-6, MDL MFCD00011256, †		POR
45930	Terbium sputtering target, 76mm dia x 3.18mm thick [7440-27-9], Tb, EINECS 231-137-6, MDL MFCD00011256, †		POR
45481	Tin shot, tear drop, Puratronic®, 99.99999% (metals basis) [7440-31-5], MDL MFCD00133862, †	5g 25g 100g	62.10 187.00 562.00

Stock #	Description	Size	\$
45584	Titanium foil, 0.127mm (0.005in) thick, annealed, 99.99+% (metals basis) [7440-32-6], MDL MFCD00011264, †	25x25mm	40.50
		50x50mm	113.00
		100x100mm	340.00
45485	Titanium wire, 1.0mm (0.04in) dia, hard, 99.99% (metals basis) [7440-32-6], ≈3.54g/m, MDL MFCD00011264, †	25cm	36.60
		100cm	110.00
		500cm	409.00
45620	Ytterbium sputtering target, 50.8mm dia x 1.59mm thick [7440-64-4], Yb, Target, m.p. 824°, b.p. 1427°, d. 6.98, EINECS 231-173-2, MDL MFCD00011286, †	each	890.00
45736	Ytterbium sputtering target, 50.8mm dia x 3.18mm thick [7440-64-4], Yb, EINECS 231-173-2, MDL MFCD00011286, †	each	995.00
46060	Ytterbium sputtering target, 76.2mm dia x 1.59mm thick [7440-64-4], Yb, EINECS 231-173-2, MDL MFCD00011286, †		POR
45928	Ytterbium sputtering target, 76.2mm dia x 3.18mm thick [7440-64-4], Yb, EINECS 231-173-2, MDL MFCD00011286, †		POR

Alloys

Stock #	Description	Size	\$
45528	Aluminum Magnesium plate, alloy 5052, 2.29mm (0.090in) thick Al:Mg:Cr; 97.25:2.5:0.25 wt%, ≈578g/30x30cm, MDL MFCD00214039, †	30x30cm	96.80
45526	Devarda's Alloy, granular [8049-11-4], Cu:Al:Zn; 50:45:5 wt%, d. 4.355, Merck 14,2941, UN3089, MDL MFCD00198176, †	100g 500g	32.30 95.70
	 R:11, S:33		
45608	Gold Beryllium pellet, 2.00mm (0.08in) dia x 10mm (0.39in) length, slightly pitted surface Au:Be; 99:1 wt%, ≈0.554g/pellet, MDL MFCD00003436, †	each 10each	85.20 785.00
	 R:49-43-48/23, S:53-4-9-20-37-45		
45489	Gold Beryllium pieces, 99.98% (metals basis) Au:Be; 99:1 wt%, Size 3.175mm-6.35mm, †	1g 5g	240.00 1000.00
45589	Misch Metal, 99.0% min. rare earth content [62379-61-7], Typically Ce:La 75:25 wt%, Pieces, UN3178, MDL MFCD00144799, †	500g 2kg	121.00 364.00
	 R:11-15, S:7/8-33-43-60		
45610	Nickel Iron Molybdenum flake, -200 mesh Ni:Fe:Mo; 81:17:2 wt%, Flakes, d. 0.29, UN3089	250g 1kg	75.00 175.00
	  R:11-40-43, S:36/37		
45492	Nitinol foil, 0.05mm (0.002in) thick, superelastic, flat annealed, pickled surface ≈0.25g/25x25mm; Af approx. 10°C, MDL MFCD02091734, †	25x25mm 50x50mm 100x100mm	53.80 183.00 540.00
	 R:40-43, S:36/37		
45514	Nitinol foil, 0.127mm (0.005in) thick, shape memory, flat annealed, pickled surface ≈0.51g/25x25mm; Af approx. 45°C, MDL MFCD02091734, †	25x25mm 50x50mm 100x100mm	55.90 188.00 645.00
	 R:40-43, S:36/37		
45578	Nitinol compression coil spring, wire diameter 1.0mm (0.039in), mean coil diameter 8.0mm (0.31in), Alloy M , †	5pcs 25pcs 100pcs	42.50 199.00 745.00
	 R:40-43, S:36/37		
45655	Nitinol tension coil spring, wire diameter 0.75mm (0.030in), mean coil diameter 5.55mm (0.22in), Alloy M , †	5pcs 25pcs 100pcs	37.50 175.00 670.00
45662	Nitinol foil, 0.018mm (0.0007in) thick, shape memory, flat annealed, pickled surface Foil, MDL MFCD02091734, †	25x25mm 50x50mm 75x100mm	130.00 380.00 1100.00
	 R:40-43, S:36/37		
45663	Nitinol foil, 0.05mm (0.002in) thick, shape memory, flat annealed, pickled surface ≈0.25g/25x25mm; Af approx. 45°C, MDL MFCD02091734, †	25x25mm 50x50mm 75x100mm	65.00 190.00 680.00
	 R:40-43, S:36/37		
45658	Nitinol wire, 1.0mm (0.039in) dia, shape memory, Alloy M, straight annealed, oxide surface ca 49.71g/10m; Af approx. 45°, †	10m	360.00
	 R:40-43, S:36/37		
45571	Platinum 30 wt% Rhodium wire, 0.33mm (0.013in) dia, ISA Type B Standard Grade Thermocouple Pt:Rh; 70:30 wt%, ≈0.75g/50cm, MDL MFCD00198186, †	50cm 2m	255.00 640.00
45516	Stainless Steel powder, APS 15 micron, Type 430-L Fe:Cr:Si:C; 82.9:16.5:0.5:0.02 wt%, †	100g 500g 2kg	37.60 96.80 296.00



Section 3 Ceramics and Materials

Alphabetical listing of Ceramics and Materials 85

This section lists other new products including ceramics, optical windows and a variety of other materials.

If you cannot find a product with the specifications you need, simply call our Speciality and Bulk Sales Department.

Stock #	Description	Size	\$
45601	Atomizer, for the microwelder	1each	425.00
45530	Boron 10, ¹⁰ B, plasma standard solution, Specpure®, ¹⁰ B 100µg/ml Matrix: 2% HNO ₃ , Liquid, b.p. 100°, d. 1.0, UN3264, †  R:34, S:23-26-36/37/39-45	50ml	156.00
45541	Boron Nitride Bar;Length (mm), 300;Width (mm), 6.4;Height (mm), 6.4 BN, Bar (Hot Pressed) 12 x 1/4 x 1/4" pc ~21g, †	1pc	302.00
45721	Boron Nitride Bar;Length (mm), 300;Width (mm), 12.7;Height (mm), 12.7 MDL MFCD00011317	1pc	400.00
45668	Boron Nitride Rectangular Plate;Length (mm), 125;Width (mm), 125;Thick (mm), 6.4 BN, Plate,HotPressed 5x5x.25" pc ~175g.	each	395.00
45598	Boron Nitride Rectangular Plate;Length (mm), 125;Width (mm), 125;Thick (mm), 12.7 BN, Plate, HotPressed 5x5x0.5" ~349g	each	660.00
45850	Boron Nitride Rod;Diameter (mm), 6.4;Length (mm), 300 [10043-11-5], Rod, m.p. 2700°, d. 2.25, EINECS 233-136-6, †	each	200.00
45912	Boron Nitride Rod;Diameter (mm), 12.7;Length (mm), 300	each	270.00
45625	Cadmium telluride Optical Window, 12.7mm (0.5in) dia x 1mm (0.04in) thick Clear Aperture 85%, Surface Finish 60/40, †	each	430.00
45685	Cadmium telluride Optical Window, 12.7mm (0.5in) dia x 2mm (0.08in) thick Clear Aperture 85%, Surface Finish 60/40, †	each	470.00
45593	Cadmium telluride Optical Window, 25.4mm (1.0in) dia x 2mm (0.08in) thick Clear Aperture 85%, Surface Finish 60/40, †	each	680.00
45637	Cadmium telluride Optical Window, 25.4mm (1.0in) dia x 4mm (0.16in) thick Clear Aperture 85%, Surface Finish 60/40, †	each	840.00
45627	Cesium iodide Optical Window, 12.7mm (0.5in) dia x 3mm (0.12in) thick Clear Aperture 85%, Surface Finish 60/40, †	each	360.00
45686	Cesium iodide Optical Window, 25.4mm (1.0in) dia x 3mm (0.12in) thick Clear Aperture 85%, Surface Finish 60/40, †	each	450.00
45647	Cesium iodide Optical Window, 50.8mm (2.0in) dia x 3mm (0.12in) thick Clear Aperture 85%, Surface Finish 60/40, †	each	1050.00
45628	Gallium arsenide Optical Window, 12.7mm (0.5in) dia x 1mm (0.04in) thick Clear Aperture 85%, Surface Finish 60/40, †	each	190.00
45667	Gallium arsenide Optical Window, 12.7mm (0.5in) dia x 2mm (0.08in) thick Clear Aperture 85%, Surface Finish 60/40, †	each	210.00
46107	Gallium arsenide Optical Window, 25.4mm (1.0in) dia x 2mm (0.08in) thick Clear Aperture 85%, Surface Finish 60/40, †	each	280.00
45636	Gallium arsenide Optical Window, 50.8mm (2.0in) dia x 2mm (0.08in) thick Clear Aperture 85%, Surface Finish 60/40, †	each	1260.00
45540	Glass Ceramic Rectangular Plate;Length (mm), 100;Width (mm), 150;Thick (mm), 6.4	each	150.00
45635	Glass Ceramic Rectangular Plate;Length (mm), 100;Width (mm), 150;Thick (mm), 9.5	each	170.00
45720	Glass Ceramic Rectangular Plate;Length (mm), 100;Width (mm), 150;Thick (mm), 12.7	each	196.00
45762	Glass Ceramic Rod;Diameter (mm), 6.4;Length (mm), 300	each	170.00
45818	Glass Ceramic Rod;Diameter (mm), 12.7;Length (mm), 300	each	200.00
45551	ICP-MS QC Standard solution 20, Specpure Matrix: 5% HNO ₃ /tr. tartaric acid, Liquid, UN3264, †	100ml	254.00
45641	ICP-MS Tuning Standard solution for 200.8, Specpure Matrix: 5% HNO ₃ , Liquid, UN3264, †	100ml	116.00
45733	Magnesium Oxide Crucible, Cylindrical, Flat Base;OD (mm), 25;Height (mm), 25	each	66.00
45854	Magnesium Oxide Crucible, Cylindrical, Flat Base;OD (mm), 32;Height (mm), 32	each	80.00
46021	Magnesium Oxide Crucible, Cylindrical, Flat Base;OD (mm), 38;Height (mm), 38	each	104.00

Stock #	Description	Size	\$
45680	Magnesium Oxide Crucible, Cylindrical, Flat Base;OD (mm), 44;Height (mm), 44	each	158.00
46108	Magnesium Oxide Crucible, Cylindrical, Flat Base;OD (mm), 51;Height (mm), 51	each	156.00
45900	Magnesium Oxide Crucible, Cylindrical, Flat Base;OD (mm), 76;Height (mm), 76	each	275.00
45749	Magnesium Oxide Rectangular Tray;Length (mm), 50;Width (mm), 50;Height (mm), 12.7	each	90.00
45954	Magnesium Oxide Rectangular Tray;Length (mm), 75;Width (mm), 75;Height (mm), 25.4	each	190.00
45634	Magnesium Oxide Rectangular Tray;Length (mm), 100;Width (mm), 25;Height (mm), 12.7	each	95.00
45827	Magnesium Oxide Rectangular Tray;Length (mm), 100;Width (mm), 50;Height (mm), 12.7	each	130.00
45791	Magnesium Oxide Rectangular Tray;Length (mm), 100;Width (mm), 100;Height (mm), 25.4	each	250.00
45865	Magnesium Oxide Rectangular Tray;Length (mm), 150;Width (mm), 75;Height (mm), 25.4	each	275.00
45699	Magnesium Oxide Rectangular Tray;Length (mm), 150;Width (mm), 100;Height (mm), 25.4	each	310.00
45555	Quartz lid for 30ml quartz crucible, fused; ID 48mm	1pc	75.00
45626	Quartz Optical Window, 12.7mm (0.5in) dia x 1mm (0.04in) thick Clear Aperture 85%, Surface Finish 40/20	each	230.00
45700	Quartz Optical Window, 12.7mm (0.5in) dia x 2mm (0.08in) thick Clear Aperture 85%, Surface Finish 40/20	each	245.00
45537	Quartz Optical Window, 25.4mm (1.0in) dia x 1mm (0.04in) thick Clear Aperture 85%, Surface Finish 40/20	each	285.00
45673	Quartz Optical Window, 25.4mm (1.0in) dia x 2mm (0.08in) thick Clear Aperture 85%, Surface Finish 40/20	each	310.00
45653	Quartz Optical Window, 25.4mm (1.0in) dia x 3mm (0.12in) thick Clear Aperture 85%, Surface Finish 40/20	each	790.00
45638	Sapphire Optical Window, 12.7mm (0.5in) dia x 1mm (0.04in) thick Clear Aperture 85%, Surface Finish 60/40	each	176.00
45560	Sapphire Optical Window, 25.4mm (1.0in) dia x 1mm (0.04in) thick Clear Aperture 85%, Surface Finish 60/40	each	198.00
45654	Sapphire Optical Window, 25.4mm (1.0in) dia x 2mm (0.08in) thick Clear Aperture 85%, Surface Finish 60/40	each	220.00
45717	Sapphire Optical Window, 50.8mm (2.0in) dia x 3mm (0.12in) thick Clear Aperture 85%, Surface Finish 60/40	each	530.00
45581	Soldering flux, liquid All purpose flux for general soldering, UN1840, MDL MFCD01868519, †  R:22-34-50/53, S:26-36/37/39-45-60-61	100ml 500ml	12.70 36.70
45568	Torch, blue, single valve	1each	180.00
45476	Tube guide assembly for Magnetic Susceptibility Balance MSB-1	1each	92.20
45486	Zinc oxide substrate, 10x10x0.5mm, polished one side, 0001 orientation ZnO, EINECS 215-171-9, MDL MFCD00011109, †  R:50/53, S:60-61	1each 5each	550.00 1926.00
45487	Zinc oxide substrate, 10x10x0.5mm, polished two sides, 0001 orientation ZnO, EINECS 215-171-9, MDL MFCD00011109, †  R:50/53, S:60-61	1each 5each	925.00 3144.00
45633	Zinc selenide Optical Window, 12.7mm (0.5in) dia x 1mm (0.04in) thick Clear Aperture 85%, Surface Finish 60/40, †	each	250.00
45666	Zinc selenide Optical Window, 12.7mm (0.5in) dia x 2mm (0.08in) thick Clear Aperture 85%, Surface Finish 60/40, †	each	260.00
45834	Zinc selenide Optical Window, 25.4mm (1.0in) dia x 1mm (0.04in) thick Clear Aperture 85%, Surface Finish 60/40, †	each	300.00
45885	Zinc selenide Optical Window, 25.4mm (1.0in) dia x 2mm (0.08in) thick Clear Aperture 85%, Surface Finish 60/40, †	each	330.00
45955	Zinc selenide Optical Window, 50.8mm (2.0in) dia x 2mm (0.08in) thick Clear Aperture 85%, Surface Finish 60/40, †	each	834.00



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