

**USAN** (The USP Dictionary of U.S. Adopted Names and International Drug Names)

<b>Subject Coverage</b>	A compilation of the U.S. Adopted Names and other drug names such as USP or NF names, and other non-proprietary names		
<b>File Type</b>	Directory, Substance		
<b>Features</b>	<a href="#">CAS Registry Numbers®</a> <input checked="" type="checkbox"/>	Page Images <input type="checkbox"/>	STN AnaVist <input type="checkbox"/>
	<a href="#">Keep &amp; Share</a> <input checked="" type="checkbox"/>	<a href="#">SLART</a> <input checked="" type="checkbox"/>	<a href="#">STN Easy</a> <input checked="" type="checkbox"/>
	Learning Database <input type="checkbox"/>	Structures <input checked="" type="checkbox"/>	STN Viewer <input type="checkbox"/>
<b>Record Content</b>	<ul style="list-style-type: none"><li>• USAN is recognized throughout the health care industry as the authoritative drug dictionary</li><li>• One or more of the following: U.S. Adopted Names, official drug names for the USP (U.S. Pharmacopeia) and NF (National Formulary), previously used official names, international and nonproprietary names, British Approved Names, Japanese Approved Names, miscellaneous older names, and trade names</li><li>• Other substance information such as molecular formula, molecular weight, pharmacological and/or therapeutic category, drug manufacturer, reference information, and structure diagram, if available</li><li>• CAS Registry Number</li></ul>		
<b>File Size</b>	10,900 records (09/11)		
<b>Coverage</b>	1953-present		
<b>Updates</b>	Reloaded annually		
<b>Language</b>	English		
<b>Database Producer</b>	The United States Pharmacopeial Convention, Inc. 12601 Twinbrook Parkway Rockville, MD 20852 USA Phone: (301) 881-0666 (International) Phone: (800) 822-8772 E-mail: <a href="mailto:webmaster@usp.org">webmaster@usp.org</a> URL: <a href="http://www.usp.org">http://www.usp.org</a>		

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**USAN**

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**Sources** USAN and the USP Dictionary of Drug Names

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**User Aids**

- Online Helps (HELP DIRECTORY lists all help messages available)
- STNGUIDE

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**Clusters**

- [CASRNS](#)
- [CHEMISTRY](#)
- [COMPANIES](#)
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**Pricing** See the [STN Price List](#) or enter HELP COST at an arrow prompt (=>).

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## Search and Display Field Codes

Fields that allow left truncation are indicated by an asterisk (\*).

Search Field Name	Search Code	Search Examples	Display Codes
Basic Index* (contains single words from the company name (CN), chemical name (CN), generic (US Adopted Name) name (CN), code designation (CN), trade name (CN), NSC number (CN), classification code (CC), and reference (RE) fields, as well as molecular formulas (MF) and CAS Registry Numbers (RN))	None (or /BI)	S BUTANOIC ACID S 3563-92-6 S ?BUTAN?/BI S C24H36O5 S ANTIDYSKINETIC	CC, CN, MF, RE, RN
Accession Number Atom Count <b>(1)</b>	/AN /ATC	S 2011:1493/AN S 20-30/ATC S ATC>30	AN Not displayed
Chemical Name (contains generic (US Adopted Name), chemical, and trade names, code designations, and NSC Numbers)	/CN	S MEVACOR/CN	CN, RE
Chemical Name Segment*	/CNS	S ?BUTYR?/CNS S ESTER/CNS	CN
Classification Code <b>(2)</b>	/CC	S ANTIHYPERLIPIDEMIC/CC S COA REDUCTASE/CC S (ANTIHYPERTENSIVE AND DIURETIC)/CC	CC
Company Name (Corporate Name) <b>(2)</b>	/CO	S MERCK/CO S MERRELL DOW/CO S TRIOLE?/CN (L) SQUIBB/CO	CN
Cross Referenced Name Element Count <b>(1)</b> Element Symbol Entry Date <b>(1)</b> (Update date)	/CR /ELC /ELS /ED (or /UP)	S MEVINOLIN/CR S 24/C AND 5/O S S/ELS AND P/ELS S ED>20110800	CR Not displayed Not displayed Not displayed
Field Availability (code and text) Linear Structural Formula Molecular Formula <b>(3)</b>	/FA /LSF /MF	S L1 AND RN/FA S C24 H36 O5/LSF S C24H36O5/MF S C24 H36 O5/MF	FA LSF MF
Molecular Weight <b>(1)</b>	/MW (or /FW)	S MW<200	MW
Number of Components <b>(1,3)</b> Periodic Group Publication Year <b>(1)</b>	/NC /PG /PY	S 1/NC S A7/PG S 2002/PY S PY>2009	Not displayed Not displayed PY
Reference Source (Compendial Reference) Trade Name	/RE /SO /TN	S BRITISH APPROVED NAME/RE S NATIONAL FORMULARY/SO S MEVACOR/TN	RE SO CN

**(1)** Numeric search field that may be searched using numeric operators or ranges.

**(2)** Search with implied (S) proximity is available in this field.

**(3)** This field contains single components.

## DISPLAY and PRINT Formats

Any combination of formats may be used to display or print answers. Multiple codes must be separated by commas or spaces, e.g., D L1 1-5 CN RN. The fields are displayed or printed in the order requested.

Hit-term highlighting is available for all fields except FA. Highlighting must be ON during SEARCH in order to use the HIT, KWIC, OCC, and QRD formats.

Format	Content	Examples
AN CC CR FA (1) LSF MF MW (FW) PY RE RN SO STF (1) STR (2) STS (1,2) TN (CO)(1)	Accession Number Classification Code Cross Referenced Name Field Availability Linear Structural Formula Molecular Formula Molecular Weight Publication Year Reference CAS Registry Number Source Flat Structure Diagram (no stereo bonds indicated) Structure Diagram (includes stereo bonds and R/S/E/Z labels when available) Stereo Structure (includes stereo bonds when available) Trade Name and Company Name	D L4 1-4 AN D L1 3 CC D CR 5-10 D 1-3,7,8 FA D LSF D MF 1-5 D L1 MW 3 D PY D RE 2 D L8 RN 1-3 D 1,4 SO D STF D 1-10 CN STR D CN STS D TN TOTAL
ALL (2) CN IDE (2)	AN, PY, CN, RN, MF, LSF, MW, RE, SO, CC, CR, STR (ALL is the default) Chemical Name (contains generic (US Adopted Name) name, chemical name, trade name and company name, code designation, and NSC number) AN, PY, CN, RN, MF, LSF, MW, STR	D 1-5 ALL D RN CN 1,3-5 D IDE
HIT (QRD) KWIC OCC (3)	Fields containing hit terms Hit terms with 20 words on either side (KeyWord-In-Context) Fields that contain hit terms and number of times they occur	D 1 5 10 HIT D KWIC NOH D OCC

(1) Custom display only.

(2) Stereo structure diagrams are available only on graphics terminals and in offline prints, or when using STN Express with *Discover!* or STN on the Web.

(3) No online display fee for this format.

## SELECT, ANALYZE, and SORT Fields

The SELECT command is used to create E-numbers containing terms taken from the specified field in an answer set.

The ANALYZE command is used to create an L-number containing terms taken from the specified field in an answer set.

The SORT command is used to rearrange the search results in either alphabetic or numeric order of the specified field(s).

Field Name	Field Code	ANALYZE/ SELECT (1)	SORT
Accession Number	AN	Y	N
CAS Registry Number	RN	Y (2)	Y
CAS Registry Number and Chemical Name	CHEM	Y (3)	N
Chemical Name	CN	Y (4) (default)	Y
Classification Code	CC	Y	Y
Company Name (Corporate Name)	CO	Y (5)	Y
Cross Referenced Name	CR	Y	N
Formula Weight	FW	N	Y
Linear Structural Formula	LSF	Y	Y
Molecular Formula	MF	Y	N
Molecular Weight	MW	N	Y
Occurrence Count of Hit Terms	OCC	N	Y
Publication Year	PY	Y	Y
Reference	RE	Y	N
Source	SO	Y	N
Trade Name	TN	Y (5)	Y

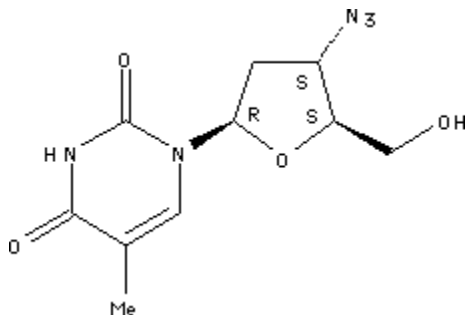
- (1) HIT may be used to restrict terms extracted to terms that match the search expression used to create the answer set, e.g., SEL HIT CN.
- (2) Appends /BI to the terms created by SELECT.
- (3) SELECTS or ANALYZES the CAS Registry Numbers, generic (US Adopted Name) name, chemical names, trade name, code designation, and NSC number and appends /BI to the terms created by SELECT.
- (4) SELECTS or ANALYZES the generic (US Adopted Name) name, chemical names, trade name, code designation, and NSC number.
- (5) SELECT HIT and ANALYZE HIT are not valid with this field.

## Sample Records

## DISPLAY ALL

Accession Number (AN): 2011:10618 USAN  
Publication Year (PY): 1987  
Generic Name (CN): Zidovudine  
OTHER NAMES:  
Chemical Name (CN): Thymidine, 3'-azido-3'-deoxy-  
Chemical Name (CN): 3'-Azido-3'-deoxythymidine  
Trade Name (CN): Retrovir (GlaxoSmithKline)  
Code Designation (CN): BW A509U; Compound S; AZT  
CAS Registry No. (RN): 30516-87-1  
Molecular Formula (MF): C<sub>10</sub> H<sub>13</sub> N<sub>5</sub> O<sub>4</sub>  
Lin. Str. Formula (LSF): C<sub>10</sub> H<sub>13</sub> N<sub>5</sub> O<sub>4</sub>  
Molecular Weight (MW): 267.24  
Reference (RE): International Nonproprietary Name; British  
Approved Name; Japanese Approved Name  
Compendial Ref. (SO): United States Pharmacopeia; USP  
Classification (CC): Antiviral  
Manufacturer Note : (Name previously used: Azidothymidine.)  
Cross Reference (CR): Azidothymidine (previously used name)

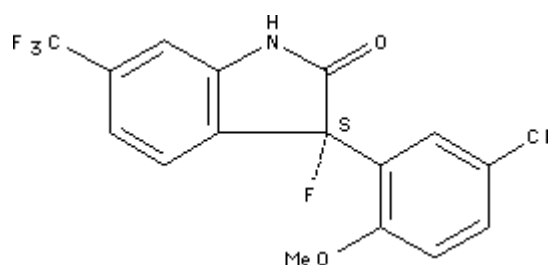
Absolute stereochemistry. Rotation (+).



## DISPLAY IDE

Accession Number (AN): 2011:4047 USAN  
Publication Year (PY): 2002  
Generic Name (CN): Flindokalner  
OTHER NAMES:  
Chemical Name (CN): 2H-Indol-2-one, 3-(5-chloro-2-methoxyphenyl)-3-fluoro-1,3-dihydro-6-(trifluoromethyl)-, (3S)-  
Chemical Name (CN): (3S)-3-(5-Chloro-2-methoxyphenyl)-3-fluoro-6-(trifluoromethyl)-1,3-dihydro-2H-indol-2-one  
Trade Name (CN): MaxiPost (Bristol-Myers Squibb)  
Code Designation (CN): BMS-204352  
CAS Registry No. (RN): 187523-35-9  
Molecular Formula (MF): C<sub>16</sub> H<sub>10</sub> Cl F<sub>4</sub> N O<sub>2</sub>  
Lin. Str. Formula (LSF): C<sub>16</sub> H<sub>10</sub> Cl F<sub>4</sub> N O<sub>2</sub>  
Molecular Weight (MW): 359.70

Absolute stereochemistry. Rotation (+).

**In North America**

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STN North America  
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Columbus, Ohio 43210-0012 U.S.A.

CAS Customer Center:  
Phone: 800-753-4227 (North America)  
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Fax: 614-447-3751  
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Internet: www.cas.org

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