

LDRUG

A training database for learning how to use the DRUG files:

DDFB Derwent Drug File Standard
 DRUGB Derwent Drug File for Subscribers

(See database summary sheet DDFU/DRUGU' for details on search and display fields, which are valid in LDRUG).

Subject Coverage	All aspects of drugs: <ul style="list-style-type: none"> • Analysis • Biochemistry • Galenics • Metabolism • Pharmacokinetics • Pharmacology • Structure-activity relationships • Synthesis • Therapeutics and adverse effects • Toxicology 		
File Type	Bibliographic, learning		
Features	Alerts (SDI)	Not available	
	CAS Registry Number® Identifiers	<input checked="" type="checkbox"/>	Page Images <input type="checkbox"/>
	Keep & Share	<input checked="" type="checkbox"/>	SLART <input type="checkbox"/>
	Learning Database	<input checked="" type="checkbox"/>	Structures <input type="checkbox"/>
Record Content	<ul style="list-style-type: none"> • Record contain bibliographic information, controlled term indexing, and structure codes (file DRUGU only) 		
File Size	<ul style="list-style-type: none"> • 500 compounds from the registry segment and about 15,000 relating literature records 		
Coverage	1983-November 1994		
Updates	Closed file		
Language	English		
Database Producer	Clarivate Friars House, 160 Black Friars Rd. London SE1 8EZ United Kingdom Copyright Holder: Clarivate		

Sources

- Medical and scientific journals and conference proceedings.
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User Aids

- Derwent Drug File Thesaurus *
 - Journal List and Selection Guidelines *
 - Product Description *
 - Work Books *
 - Online Helps (HELP DIRECTORY lists all help messages available)
 - STNGUIDE
- * available from producer
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Cluster

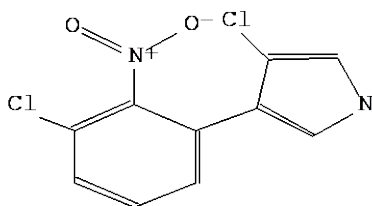
- LEARNING
- STN Database Cluster information:
<http://www.stn-international.com/en/customersupport/customer-support#cluster+%7C+subjects+%7C+features>
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Related Databases

- DDFU
 - DRUGU
- contain data from 1983 to present
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Sample Records**DISPLAY ALL STR from REGISTRY SEGMENT**

AN 20825 LDRUG
 DDRN PYRROLNIT
 DDN PYRROLNITRIN
 RN 1018-71-9
 CT ANTIBIOTICS; FUNGICIDES
 SS PYRROLE; CHLORINE; ARYLCHLORIDE; NITROARENE; BH-LINKED-CC
 MPC 02& *G; 02- *G; 03& *G; 032 *G; 06& *G; 067 *G; 071 *G; 097 *G; 10- *G;
 104 *G; 105 *G; 131 *G; 163 *G; 17& *G; 17- *G; 171 *G; 174 *G

**DISPLAY ALL from LITERATURE SEGMENT**

AN 1986-21902 LDRUG P A
 TI On the Pharmaceutical and Biopharmaceutical Assessment of Acetylamino nitropropoxybenzene (Falimint). Part 6. Quantitive Composition of the Metabolites.
 AU Gerlach K; Metzner J; Fuerst W
 LO Halle, Germany, East
 SO Pharmazie (40, Number 8, 564-65, 1985) 2 Fig. 5 Tab. 2 Reference
 CODEN: PHARAT ISSN: 0031-7144
 AV Sekt. Pharmazie der Martin-Luther-University Halle-Wittenberg, DDR-4020 Halle/Saale, Weinbergweg 15, DDR
 LA German
 DT Journal
 AB The metabolism of Falimint following p.o. administration to human subjects was studied. Polarographic analysis of the urine showed that most of the drug was eliminated within 6 hr, the main metabolite being falimintic acid (3-(2-acetamido 4-nitrophenoxy) propionic acid). Other metabolites include free and conjugated 2-acetamido-4-nitrophenol (AANP), 2-aminonitrophenol (ANP), 3-(2-amino 4-nitrophenoxy) propionic acid (M7) and 2-amino 4-nitropropoxybenzene (Ultrasweet). Variations in the relative amounts of the metabolites over the test period were discussed.
 SH P Pharmacology
 A Analysis
 CC 5 Analysis
 8 Pharmacokinetics
 33 Respiratory
 CT QUANT. *FT; DET. *FT; ANALYSIS *FT; POLAROGRAPHY *FT; ELIMINATION *FT; CONC. *FT; URINE *FT; HUMAN *FT; PHARMACOKINETICS *FT; METABOLITE *FT; ELECTROCHEM. *FT
 [01] FALIMINT *OC; FALIMINT *DM; P.O. *FT; ANTITUSSIVES *FT; FALIMINT *RN; OC *FT; DM *FT
 [02] FALIMINTATE *OC; FALIMINTATE *DM; PHENOL-ETHER *FT; NITROARENE *FT; ARYLAMINE *FT; AMINOACID *FT; C-AMIDE *FT; OC *FT; DM *FT; FALIMINTA

LDRUG

*RN

[03] ULTRASWEET *OC; ULTRASWEET *DM; ULTRASWEE *RN; OC *FT; DM *FT

[04] NITROARENE *FT; ARYLAMINE *FT; PHENOL *FT; C-AMIDE *FT; CONJUGATE *FT;
PHENOL-ETHER *FT; C-ACID *FT; OC *FT; DM *FT

FA AB; LA; CT; MPC

FS Literature

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