

Eagle Pharmaceuticals Inc

CORTELLIS COMPANY DETAILED PIPELINE REPORT

A comprehensive coverage of the the company's drug pipeline portfolio including detailed product records.

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THOMSON REUTERS

3 Times Square
New York, New York 10036
United States

Tel: +1 646 223 4000

thomsonreuters.com

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GLOSSARY

Number of Drugs in Active Development

Number of drugs associated with the company or subsidiary that are currently in active development, i.e. the development status for the drug(s) is one of the following: Discovery, Clinical, Phase I, Phase II, Phase III, Pre-registration, Registered, Launched, or Suspended.

Number of Inactive Drugs

Number of drugs associated with the company or subsidiary that are currently classified as inactive, i.e. where the development status for the drug(s) is one of the following: No Development Reported, Discontinued, or Withdrawn.

Number of Patents as Owner

Number of patents associated with the company where the company is listed as owner; i.e. the relationship type (or way the patent refers to the company) is: Patent Assignee/Owner, Patent owner (not assignee), Licensee for development and marketing, Licensee – marketing only (Distributor), Patent assignee of family member, Inferred assignee.

Number of Patents as Third Party

Number of patents associated with the company where the company is listed as third party; i.e. the relationship type (or way the patent refers to the company) is: Patent assignee (not owner), Ex-Licensee for development and marketing, Ex-Licensee marketing only (Distributor), Customer of technology, Ex-Customer of technology, Patent opponent or infringer, Affiliate organization of inventor, Owner of underlying technology.

Patents summary table

This table represents a summary of the core patent coverage for this company covering Therapeutic EP, US and WO patents since 1990 only.

Number of Deals

A count of deals where the company or one of its subsidiaries is the primary company.

Key Indications

Displays top ten key indications for the company and its subsidiaries based on frequency (indications occurring with high and identical frequency are always included, and this may result in more than ten Key Indications being listed). Includes both indications associated with patents where the company is patent owner and indications associated with drugs in active development. A drug is classified as 'active' if it features on a row (or rows) in the current development status table where the status is one of the following: Discovery, Clinical, Phase I, Phase II, Phase III, Pre-registration, Registered, Launched, or Suspended.

Key Target-based Actions

Displays top ten key target-based actions for the company and its subsidiaries based on frequency (actions occurring with high and identical frequency are always included, and this may result in more than ten Key Target-based Actions being listed). Includes both target-based actions associated with patents where the company patent owner and target-based actions associated with drugs in active development. A drug is classified as 'active' if it features on a row (or rows) in the current development status table where the status is one of the following: Discovery, Clinical, Phase I, Phase II, Phase III, Pre-registration, Registered, Launched, or Suspended. A target-based action is one that is associated with a target.

Key Technologies

Displays top ten key technologies for the company and its subsidiaries based on frequency (technologies occurring with high and identical frequency are always included, and this may result in more than ten Key Technologies being listed). Includes both key technologies associated with patents where the company relationship is patent owner and key technologies associated with drugs in active development. A drug is classified as 'active' if it features on a row (or rows) in the current development status table where the status is one of the following: Discovery, Clinical, Phase I, Phase II, Phase III, Pre-registration, Registered, Launched, or Suspended.

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Eagle Pharmaceuticals Inc

COMPANY OVERVIEW

| | |
|---------------------------------------|--|
| Company Name | Eagle Pharmaceuticals Inc |
| Parent Company Name | Eagle Pharmaceuticals Inc |
| Website | http://www.eagleus.com/ |
| Country | US |
| Number of Drugs in Active Development | 5 |
| Number of Inactive Drugs | 3 |
| Number of Patents as Owner | 11 |
| Number of Patents as Third Party | 0 |
| Number of Deals | 9 |
| Key Indications | Cancer,Non-Hodgkin lymphoma,Chronic lymphocytic leukemia,Hyperthermia,Mesothelioma,Thrombosis,Malignant hyperthermia,Metastatic non small cell lung cancer,B-cell lymphoma,Acute myelogenous leukemia,Bacterial endocarditis,Bacterial infection,Bacterial skin infection,Congestive heart failure,Enterococcus faecalis infection,MRSA infection,Myelodysplastic syndrome,Non-small-cell lung cancer,Renal insufficiency,Sepsis,Staphylococcus aureus infection,Streptococcus agalactiae infection,Streptococcus infection,Streptococcus pyogenes infection |
| Key Target-based Actions | Factor IIa antagonist,PARP modulator,DHFR inhibitor,Thymidylate synthase inhibitor,Transferase inhibitor,DNA methyltransferase inhibitor,Ryanodine receptor antagonist |
| Key Technologies | Intravenous formulation,Small molecule therapeutic,Infusion,Peptide,Biological therapeutic,Nanoparticle formulation injectable,Suspension,Formulation preservation,Liquid formulation,Injectable formulation |

COMPANY PROFILE

SUMMARY

Eagle Pharmaceuticals Inc is a specialty pharmaceutical company, founded in 2007, which develops improved formulations of injectable products.

LICENSING AGREEMENTS

In September 2009, The Medicines Company licensed rights in the US and Canada to a ready-to-use formulation of argatroban from Eagle Pharmaceuticals. The formulation was under review by the FDA. Eagle would receive \$5 million up front in cash, and The Medicines Company would make a \$2 million equity investment in Eagle. Eagle could also receive milestones and royalties.

FINANCIAL

By May 2014, a one-for-6.41 reverse stock split of the company's common stock, and all outstanding shares of preferred stock converted into 7,487,928 million shares of common stock, had been approved by the company's board of directors. And also, all series C preferred stock warrants outstanding have been exercised for 34,074 shares of common stock.

In February 2014, the company planned to issue 3.35 million shares of common stock through an initial public offering at a price of \$15 per share and underwriters were granted 30-day option to purchase additional 502,500 shares of common stock. The Offering was expected to close on February 18, 2014. At that time, the shares were expected to be traded under the ticker symbol "EGRX".

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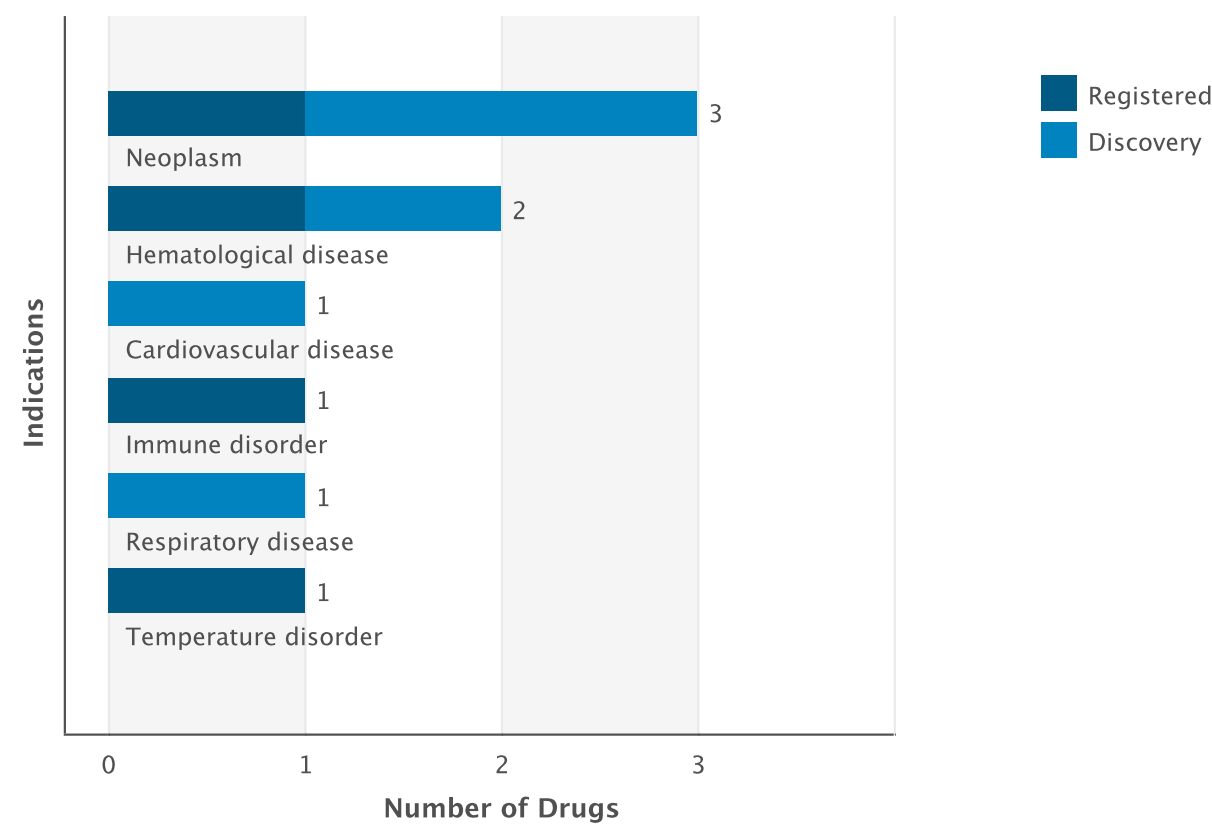


PRODUCT PORTFOLIO SUMMARY

DRUGS

Drugs by Indication

Active Drugs by Indication Chart



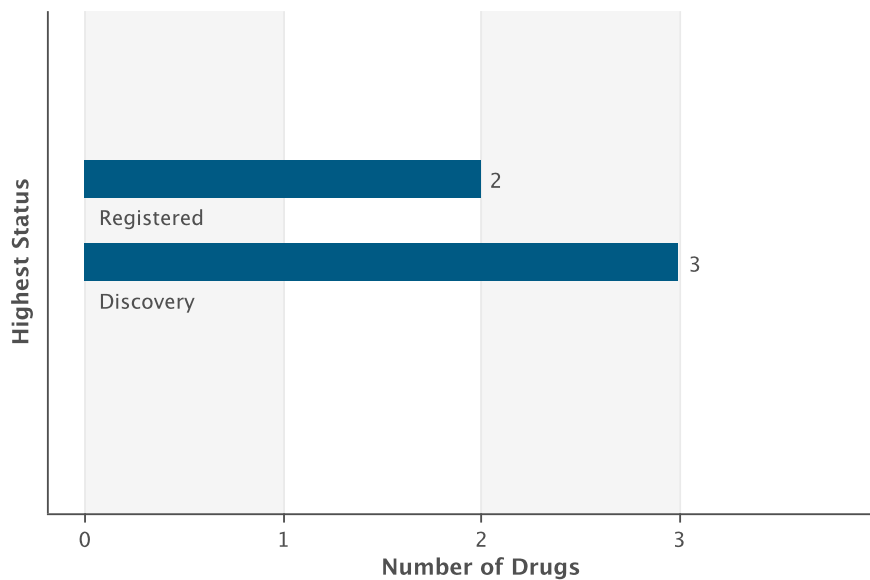
Drugs by Indication Table

| Indication | Active | Inactive | Total |
|------------------------|--------|----------|-------|
| Neoplasm | 3 | 1 | 4 |
| Hematological disease | 2 | 1 | 3 |
| Respiratory disease | 1 | 1 | 2 |
| Cardiovascular disease | 1 | 0 | 1 |
| Immune disorder | 1 | 0 | 1 |
| Temperature disorder | 1 | 0 | 1 |
| Infectious disease | 0 | 1 | 1 |

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Drugs by Highest Status

Active Drugs by Highest Status Chart



Drugs by Highest Status Table

| Development Status | Number of Drugs |
|-------------------------|-----------------|
| Registered | 2 |
| Discovery | 3 |
| Discontinued | 1 |
| No Development Reported | 2 |

DEALS

| Deal Type | Principal | | Partner | | Total |
|--|-----------|----------|---------|----------|-------|
| | Active | Inactive | Active | Inactive | |
| Patent - Exclusive Rights | 0 | 0 | 3 | 0 | 3 |
| Drug - Development/Commercialization License | 1 | 0 | 2 | 0 | 3 |
| Drug - Manufacturing/Supply | 0 | 0 | 1 | 0 | 1 |
| Technology - Delivery/Formulation | 0 | 0 | 1 | 0 | 1 |

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CLINICAL TRIALS

Trials by Condition Studied

| Condition Studied | Ongoing | All |
|--------------------------|---------|-----|
| Gastrointestinal disease | 0 | 1 |
| Hematological disease | 0 | 1 |
| Neoplasm | 0 | 1 |
| Inflammatory disease | 0 | 1 |
| Immune disorder | 0 | 1 |

Trials by Phase

| Phase | Ongoing | All |
|---------|---------|-----|
| Phase 3 | 0 | 1 |
| Phase 1 | 0 | 1 |

Phase Definitions

Phase 3 Clinical

Includes Phase 3, Phase 3b, Phase 3a, Phase 2/3 (where enrolment count is 300 or over)

Phase 2 Clinical

Includes Phase 2, Phase 2a, Phase 2b, Phase 1/2 (where enrolment count is 100 or over), Phase 2/3 (where enrolment count is under 300 or not specified)

Phase 1 Clinical

Includes Phase 1, Phase 1a, Phase 1, Phase 1/2 (where enrolment count is under 100 or not specified), Phase 0

PATENTS *

| Indication | As Owner | As Third Party | Total |
|--------------------------|----------|----------------|-------|
| Cardiovascular disease | 5 | 0 | 5 |
| Endocrine disease | 1 | 0 | 1 |
| Gastrointestinal disease | 1 | 0 | 1 |
| Genitourinary disease | 2 | 0 | 2 |
| Hematological disease | 5 | 0 | 5 |
| Immune disorder | 2 | 0 | 2 |
| Neoplasm | 6 | 0 | 6 |
| Neurological disease | 2 | 0 | 2 |
| Respiratory disease | 2 | 0 | 2 |
| Infectious disease | 1 | 0 | 1 |

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| | | | |
|---------------------------|---|---|---|
| Injury | 2 | 0 | 2 |
| Inflammatory disease | 1 | 0 | 1 |
| Gynecology and obstetrics | 1 | 0 | 1 |
| Temperature disorder | 2 | 0 | 2 |
| Dermatological disease | 1 | 0 | 1 |

* This table represents a summary of the core patent coverage for this company covering Therapeutic EP, US and WO patents since 1990 only.

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PRODUCT PORTFOLIO DRUG PIPELINE DETAIL

PLEASE NOTE: Highest status refers to highest development of that drug for one of the active companies

dantrolene sodium (heat stroke/malignant hyperthermia), Eagle Pharmaceuticals

dantrolene sodium (heat stroke/malignant hyperthermia), Eagle Pharmaceuticals SNAPSHOT

| | |
|----------------------|---|
| Drug Name | dantrolene sodium (heat stroke/malignant hyperthermia), Eagle Pharmaceuticals |
| Key Synonyms | dantrolene;dantrolene sodium;Ryanodex |
| Originator Company | Eagle Pharmaceuticals Inc |
| Active Companies | Eagle Pharmaceuticals Inc |
| Inactive Companies | |
| Highest Status | Registered |
| Active Indications | Malignant hyperthermia;Hyperthermia |
| Target-based Actions | |
| Other Actions | Unspecified drug target |
| Technologies | Nanoparticle formulation injectable;Small molecule therapeutic;Intravenous formulation;Suspension |
| Last Change Date | 09-Jan-2015 |

dantrolene sodium (heat stroke/malignant hyperthermia), Eagle Pharmaceuticals DEVELOPMENT PROFILE

SUMMARY

Eagle Pharmaceuticals has developed Ryanodex, dantrolene sodium, as a nanoparticle suspension for iv injection, for the potential treatment of malignant hyperthermia and also developing the drug as EP-4104, for the potential treatment of heat stroke,. In July 2014, the FDA had approved the drug for malignant hyperthermia and the company was planning to ship the drug in the very 'near future'. By March 2014, a phase I study for EP-4104 had been completed. In December 2014, a pivotal trial of Ryanodex for exertional heat stroke was expected to be initiated by the end of 2015. In August 2014, the company was planning to file for approval of Ryanodex with the EMA by mid-2015 .

REGULATORY INFORMATIONTHE US

In January 2014, an NDA was filed to the US FDA for the treatment of malignant hyperthermia. In March 2014, the US FDA accepted the NDA and granted a Priority Review classification. At that time, the PDUFA date was given as July 22, 2014 and the company intended to commercialize dantrolene after approval and retain exclusive marketing rights in the US. In July 2014, the FDA had approved the drug for malignant hyperthermia; at that time, the drug was expected to be available in August 2014 and the company was planning to ship the drug in the very 'near future'.

In September 2012, the US FDA granted dantrolene sodium Orphan Drug designation for the treatment of heat stroke. In August 2013, the FDA granted dantrolene sodium suspension for injection Orphan designation for the treatment of malignant hyperthermia syndrome. In July 2014, the drug was expected to receive the seven year Orphan Drug market exclusivity in 4 to 6 weeks.

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EUROPE

In October 2014, the EMA's COMP recommended granting dantrolene sodium Orphan designation for the treatment of malignant hyperthermia ; in November 2014, the Orphan Drug status was granted.

In August 2014, the company was planning to file for approval of Ryanodex with the EMA by mid-2015 .

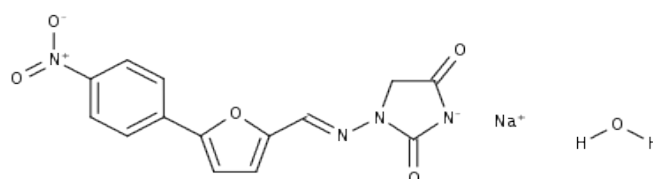
dantrolene sodium (heat stroke/malignant hyperthermia), Eagle Pharmaceuticals DEVELOPMENT STATUS

CURRENT DEVELOPMENT STATUS

| Company | Indication | Country | Development Status | Date |
|---------------------------|------------------------|---------|--------------------|-------------|
| Eagle Pharmaceuticals Inc | Malignant hyperthermia | US | Registered | 23-Jul-2014 |
| Eagle Pharmaceuticals Inc | Hyperthermia | US | Discovery | 25-Sep-2012 |
| Eagle Pharmaceuticals Inc | Malignant hyperthermia | Europe | Discovery | 11-Aug-2014 |

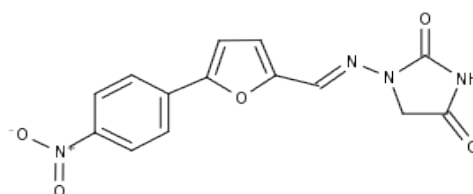
dantrolene sodium (heat stroke/malignant hyperthermia), Eagle Pharmaceuticals CHEMICAL STRUCTURES

| CAS Registry Number: | Confidence Level: |
|----------------------|-------------------|
| 24868-20-0 | 2 |



| Name | Type |
|-------------------|------|
| dantrolene sodium | USAN |

| CAS Registry Number: | Confidence Level: |
|----------------------|-------------------|
| 7261-97-4 | 1 |



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| Name | Type |
|------------|-----------------|
| dantrolene | BANN; INN; USAN |
| Ryanodex | Trade Name |

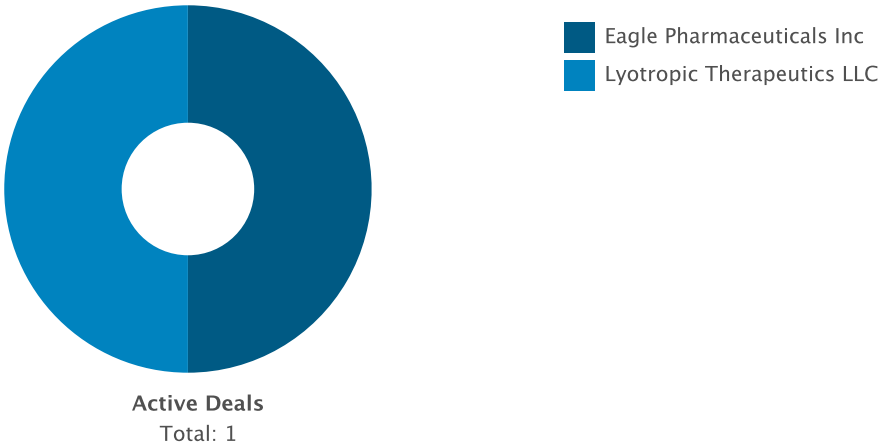
dantrolene sodium (heat stroke/malignant hyperthermia), Eagle Pharmaceuticals DRUG NAMES

| Names | Type |
|---|-----------------|
| EP-4104 | Research Code |
| dantrolene sodium | USAN |
| dantrolene | BANN, INN, USAN |
| dantrolene sodium (heat stroke/malignant hyperthermia), Eagle Pharmaceuticals | |
| Ryanodex | Trade Name |

dantrolene sodium (heat stroke/malignant hyperthermia), Eagle Pharmaceuticals DEALS AND PATENTS

DEALS

Deals by Parent Company Chart



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Deals by Parent Company Table

| Company Name | Principal | | Partner | | Total |
|----------------------------|-----------|----------|---------|----------|-------|
| | Active | Inactive | Active | Inactive | |
| Lyotropic Therapeutics LLC | 1 | 0 | 0 | 0 | 1 |
| Eagle Pharmaceuticals Inc | 0 | 0 | 1 | 0 | 1 |

Deals by Type Chart

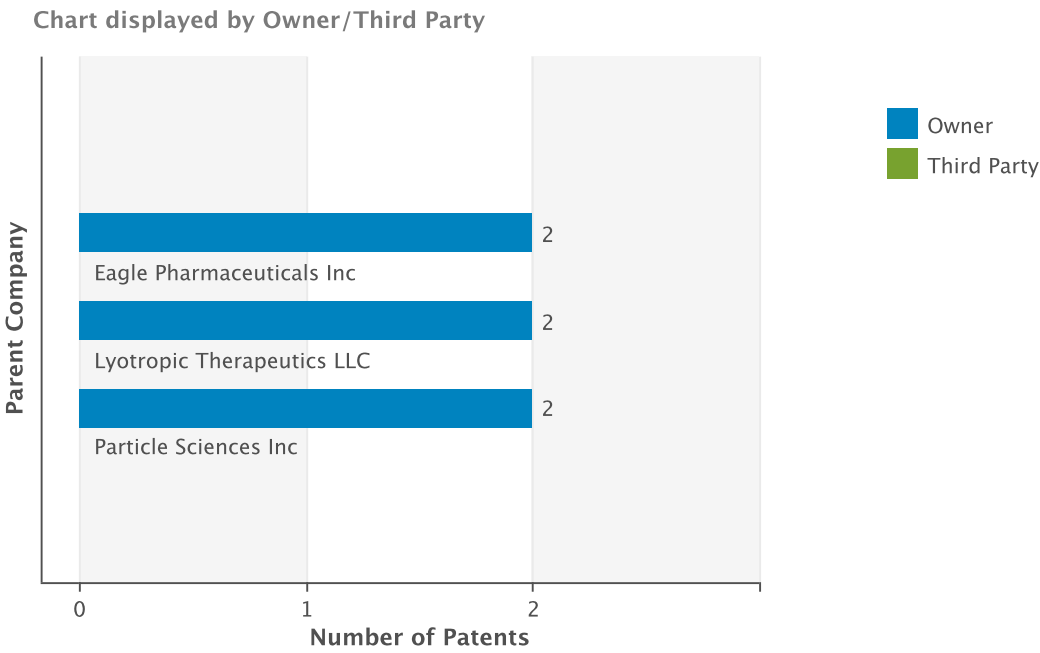


Deals by Type Table

| Deal Type | Active | Inactive | Total |
|---------------------------|--------|----------|-------|
| Patent - Exclusive Rights | 1 | 0 | 1 |

PATENTS

Patents by Parent Company Chart

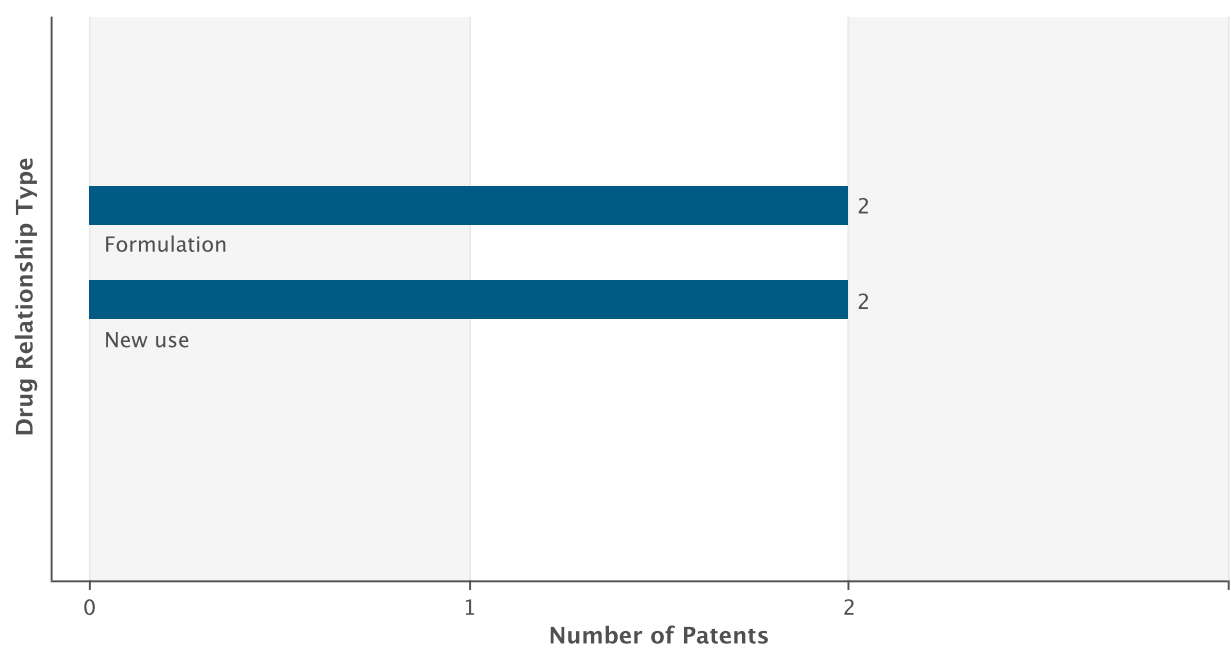


Patents by Parent Company Table

| Company Name | As Owner | As Third Party | Total |
|----------------------------|----------|----------------|-------|
| Particle Sciences Inc | 2 | 0 | 2 |
| Lyotropic Therapeutics LLC | 2 | 0 | 2 |
| Eagle Pharmaceuticals Inc | 2 | 0 | 2 |

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Patents by Drug Relationship Type Chart



Patents by Drug Relationship Type Table

| Drug Relationship | Total |
|-------------------|-------|
| Formulation | 2 |
| New use | 2 |

ready-to-dilute bendamustine hydrochloride (iv, chronic lymphocytic leukemia/non-Hodgkin's lymphoma), Eagle Pharmaceuticals

ready-to-dilute bendamustine hydrochloride (iv, chronic lymphocytic leukemia/non-Hodgkin's lymphoma), Eagle Pharmaceuticals **SNAPSHOT**

| | |
|-----------------------------|---|
| Drug Name | ready-to-dilute bendamustine hydrochloride (iv, chronic lymphocytic leukemia/non-Hodgkin's lymphoma), Eagle Pharmaceuticals |
| Key Synonyms | bendamustine |
| Originator Company | Eagle Pharmaceuticals Inc |
| Active Companies | Eagle Pharmaceuticals Inc |
| Inactive Companies | |
| Highest Status | Registered |
| Active Indications | Chronic lymphocytic leukemia;Non-Hodgkin lymphoma |
| Target-based Actions | PARP modulator |
| Other Actions | Anticancer alkylating agent |
| Technologies | Small molecule therapeutic;Intravenous formulation;Infusion |
| Last Change Date | 18-Dec-2014 |

ready-to-dilute bendamustine hydrochloride (iv, chronic lymphocytic leukemia/non-Hodgkin's lymphoma), Eagle Pharmaceuticals **DEVELOPMENT PROFILE**

SUMMARY

Eagle Pharmaceuticals has developed a ready-to-dilute (RTD) 500-ml iv formulation of bendamustine hydrochloride, an alkylating agent and PARP modulator, as EP-3101, for the potential treatment of chronic lymphocytic leukemia (CLL) and non-Hodgkin's lymphoma (NHL); in July 2014, the FDA granted tentative approval for the RTD formulation of bendamustine for indolent B-cell NHL; final approval is subject to the resolution of ongoing patent litigation between Eagle and Teva and the resolution or expiry of certain Orphan Drug exclusivities held by Teva .

The company is also developing a 50-ml iv formulation of bendamustine hydrochloride with short infusion time, as EP-3102, for the potential treatment of CLL and NHL. In November 2013, a phase I trial began; in November 2014, data were reported. At that time, the company planned a pre-NDA meeting with the FDA in mid-December 2014.

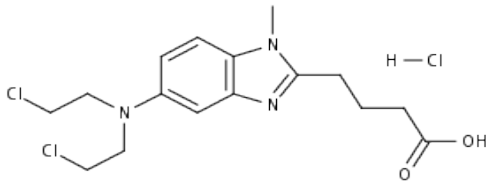
ready-to-dilute bendamustine hydrochloride (iv, chronic lymphocytic leukemia/non-Hodgkin's lymphoma), Eagle Pharmaceuticals **DEVELOPMENT STATUS**

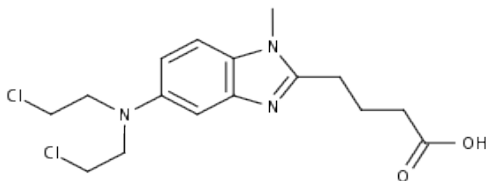
CURRENT DEVELOPMENT STATUS

| Company | Indication | Country | Development Status | Date |
|---------------------------|------------------------------|---------|--------------------|-------------|
| Eagle Pharmaceuticals Inc | Non-Hodgkin lymphoma | US | Registered | 02-Jul-2014 |
| Eagle Pharmaceuticals Inc | Chronic lymphocytic leukemia | US | Pre-registration | 06-Sep-2013 |

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ready-to-dilute bendamustine hydrochloride (iv, chronic lymphocytic leukemia/non-Hodgkin's lymphoma), Eagle Pharmaceuticals **CHEMICAL STRUCTURES**

| | |
|--|-------------------|
| CAS Registry Number: | Confidence Level: |
| 3543-75-7 | 2 |
|  | |
| Name | Type |
| Treakisym | Trade Name |
| EP-3102 | Research Code |
| EP-3101 | Research Code |
| bendamustine hydrochloride | |

| | |
|--|-------------------|
| CAS Registry Number: | Confidence Level: |
| 16506-27-7 | 1 |
|  | |
| Name | Type |
| bendamustine | INN |
| Treanda | Trade Name |
| Levact | Trade Name |
| SP-1031C | Research Code |
| SDX-105 | Research Code |

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ready-to-dilute bendamustine hydrochloride (iv, chronic lymphocytic leukemia/non-Hodgkin's lymphoma), Eagle Pharmaceuticals DRUG NAMES

| Names | Type |
|--|---------------|
| bendamustine hydrochloride | |
| EP-3102 | Research Code |
| ready-to-dilute bendamustine hydrochloride (iv, chronic lymphocytic leukemia/non-Hodgkin's lymphoma), Eagle Pharmaceuticals bendamustine | INN |
| EP-3101 | Research Code |

ready-to-dilute bendamustine hydrochloride (iv, chronic lymphocytic leukemia/non-Hodgkin's lymphoma), Eagle Pharmaceuticals CLINICAL TRIALS

Trials by Phase and Condition Studied

| Phase 4 Clinical | | Phase 3 Clinical | | Phase 2 Clinical | | Phase 1 Clinical | | Phase Unspecified | | Total | |
|------------------------|-----|------------------|-----|------------------|-----|------------------|-----|-------------------|-----|----------|-----|
| On-going | All | On-going | All | On-going | All | On-going | All | On-going | All | On-going | All |
| Solid tumor | | | | | | | | | | | |
| 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 1 |
| Hematological neoplasm | | | | | | | | | | | |
| 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 1 |

Total Trials by Phase and Status

| Phase 4 Clinical | | Phase 3 Clinical | | Phase 2 Clinical | | Phase 1 Clinical | | Phase Unspecified | | Total | |
|---------------------------|-----|------------------|-----|------------------|-----|------------------|-----|-------------------|-----|----------|-----|
| On-going | All | On-going | All | On-going | All | On-going | All | On-going | All | On-going | All |
| Total by Phase and Status | | | | | | | | | | | |
| 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 1 |

Phase Definitions

Phase 3 Clinical

Includes Phase 3, Phase 3b, Phase 3a, Phase 2/3 (where enrolment count is 300 or over)

Phase 2 Clinical

Includes Phase 2, Phase 2a, Phase 2b, Phase 1/2 (where enrolment count is 100 or over), Phase 2/3 (where enrolment count is under 300 or not specified)

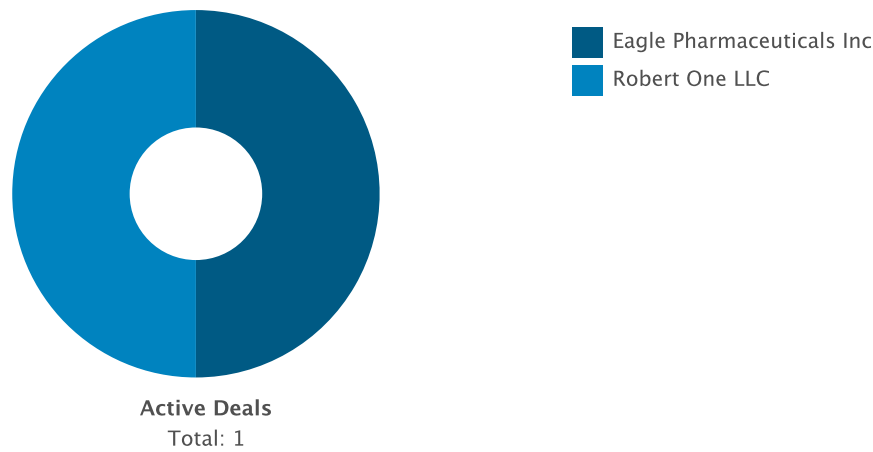
Phase 1 Clinical

Includes Phase 1, Phase 1a, Phase 1, Phase 1/2 (where enrolment count is under 100 or not specified), Phase 0

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DEALS

Deals by Parent Company Chart



Deals by Parent Company Table

| Company Name | Principal | | Partner | | Total |
|---------------------------|-----------|----------|---------|----------|-------|
| | Active | Inactive | Active | Inactive | |
| Eagle Pharmaceuticals Inc | 0 | 0 | 1 | 0 | 1 |
| Robert One LLC | 1 | 0 | 0 | 0 | 1 |

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Deals by Type Chart



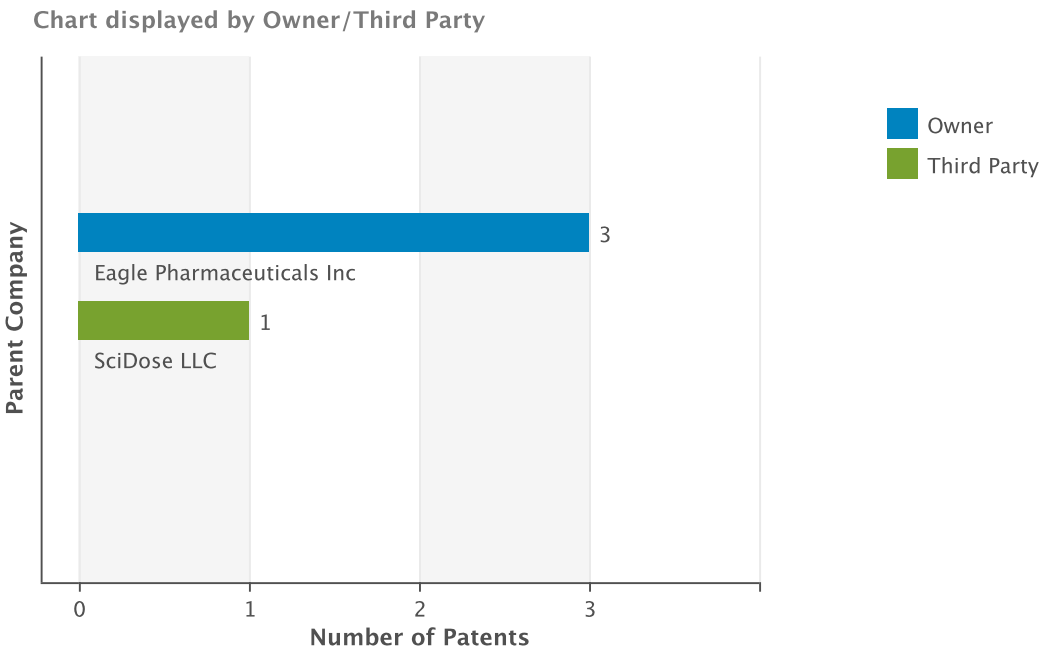
Deals by Type Table

| Deal Type | Active | Inactive | Total |
|---------------------------|--------|----------|-------|
| Patent - Exclusive Rights | 1 | 0 | 1 |

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PATENTS

Patents by Parent Company Chart

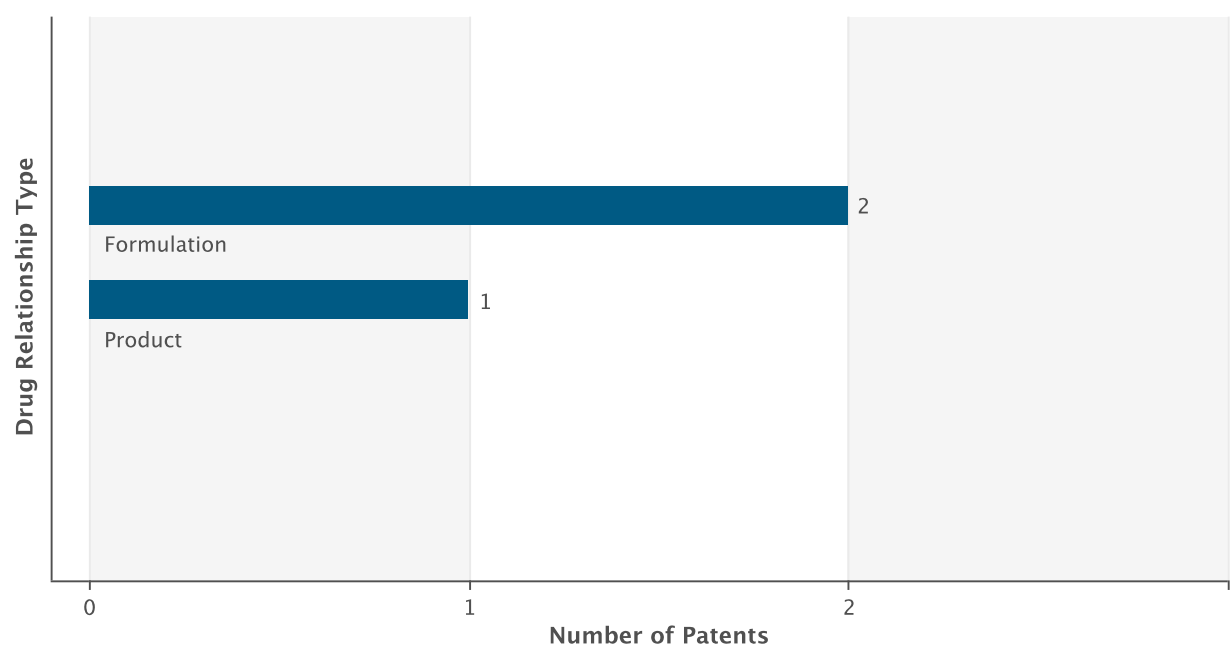


Patents by Parent Company Table

| Company Name | As Owner | As Third Party | Total |
|---------------------------|----------|----------------|-------|
| Eagle Pharmaceuticals Inc | 3 | 0 | 3 |
| SciDose LLC | 0 | 1 | 1 |

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Patents by Drug Relationship Type Chart



Patents by Drug Relationship Type Table

| Drug Relationship | Total |
|-------------------|-------|
| Formulation | 2 |
| Product | 1 |

ready-to-use bivalirudin (iv, thrombosis in percutaneous transluminal angioplasty), Eagle Pharmaceuticals

ready-to-use bivalirudin (iv, thrombosis in percutaneous transluminal angioplasty), Eagle Pharmaceuticals SNAPSHOT

| | |
|----------------------|---|
| Drug Name | ready-to-use bivalirudin (iv, thrombosis in percutaneous transluminal angioplasty), Eagle Pharmaceuticals |
| Key Synonyms | bivalirudin |
| Originator Company | Eagle Pharmaceuticals Inc |
| Active Companies | Eagle Pharmaceuticals Inc |
| Inactive Companies | |
| Highest Status | Discovery |
| Active Indications | Thrombosis |
| Target-based Actions | Factor IIa antagonist |
| Other Actions | Coagulation inhibitor;Hirudin modulator |
| Technologies | Biological therapeutic;Intravenous formulation;Peptide |
| Last Change Date | 12-Aug-2014 |

ready-to-use bivalirudin (iv, thrombosis in percutaneous transluminal angioplasty), Eagle Pharmaceuticals DEVELOPMENT PROFILE

SUMMARY

Eagle Pharmaceuticals is investigating EP-6101, a ready-to-use (RTU) formulation of bivalirudin, an iv anticoagulant thrombin inhibitor and synthetic 20 amino acid hirudin derivative, for the potential prevention of thrombosis in percutaneous transluminal angioplasty. In March 2014, the program was listed as being in formulation and toxicology studies. In August 2014, the company was planning to submit the NDA in the first half of 2015.

ready-to-use bivalirudin (iv, thrombosis in percutaneous transluminal angioplasty), Eagle Pharmaceuticals DEVELOPMENT STATUS

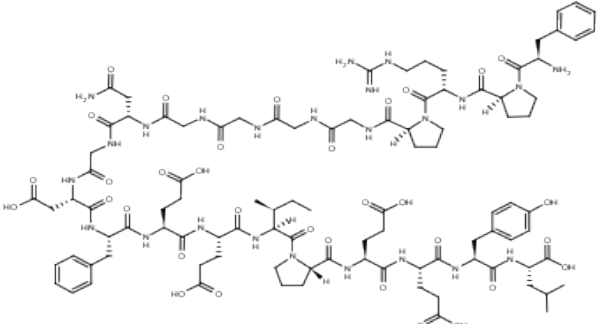
CURRENT DEVELOPMENT STATUS

| Company | Indication | Country | Development Status | Date |
|---------------------------|------------|---------|--------------------|-------------|
| Eagle Pharmaceuticals Inc | Thrombosis | US | Discovery | 30-Nov-2013 |

ready-to-use bivalirudin (iv, thrombosis in percutaneous transluminal angioplasty), Eagle Pharmaceuticals CHEMICAL STRUCTURES

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| | |
|--|-------------------|
| CAS Registry Number: | Confidence Level: |
| 128270-60-0 | 1 |
|  | |
| Name | Type |
| bivalirudin | INN; USAN |
| Hirulog | Trade Name |
| Angiomax | Trade Name |
| EP-6101 | Research Code |
| Hirulog-8 | |

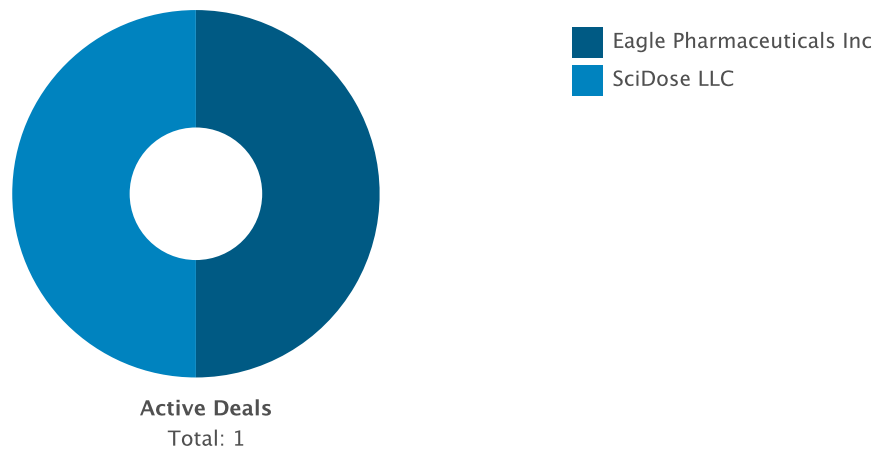
ready-to-use bivalirudin (iv, thrombosis in percutaneous transluminal angioplasty), Eagle Pharmaceuticals DRUG NAMES

| | |
|---|---------------|
| Names | Type |
| EP-6101 | Research Code |
| ready-to-use bivalirudin (iv, thrombosis in percutaneous transluminal angioplasty), Eagle Pharmaceuticals | |
| bivalirudin | INN, USAN |

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DEALS

Deals by Parent Company Chart



Deals by Parent Company Table

| Company Name | Principal | | Partner | | Total |
|---------------------------|-----------|----------|---------|----------|-------|
| | Active | Inactive | Active | Inactive | |
| Eagle Pharmaceuticals Inc | 0 | 0 | 1 | 0 | 1 |
| SciDose LLC | 1 | 0 | 0 | 0 | 1 |

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Deals by Type Chart



Deals by Type Table

| Deal Type | Active | Inactive | Total |
|---------------------------|--------|----------|-------|
| Patent - Exclusive Rights | 1 | 0 | 1 |

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undisclosed compounds (cancer), SciDose/Eagle Pharmaceuticals

undisclosed compounds (cancer), SciDose/Eagle Pharmaceuticals SNAPSHOT

| | |
|----------------------|---|
| Drug Name | undisclosed compounds (cancer), SciDose/Eagle Pharmaceuticals |
| Key Synonyms | |
| Originator Company | SciDose LLC |
| Active Companies | Eagle Pharmaceuticals Inc;SciDose LLC |
| Inactive Companies | |
| Highest Status | Discovery |
| Active Indications | Cancer |
| Target-based Actions | |
| Other Actions | Anticancer;Unspecified drug target |
| Technologies | Small molecule therapeutic |
| Last Change Date | 04-Jul-2014 |

undisclosed compounds (cancer), SciDose/Eagle Pharmaceuticals DEVELOPMENT PROFILE

SUMMARY

SciDose in collaboration with Eagle Pharmaceuticals, is investigating undisclosed compounds, for the potential treatment of cancer. In March 2011, development of nine drugs was underway. In July 2014, development of the program was presumed to be ongoing.

undisclosed compounds (cancer), SciDose/Eagle Pharmaceuticals DEVELOPMENT STATUS

CURRENT DEVELOPMENT STATUS

| Company | Indication | Country | Development Status | Date |
|---------------------------|------------|---------|--------------------|-------------|
| Eagle Pharmaceuticals Inc | Cancer | US | Discovery | 04-Mar-2011 |
| SciDose LLC | Cancer | US | Discovery | 04-Mar-2011 |

undisclosed compounds (cancer), SciDose/Eagle Pharmaceuticals DRUG NAMES

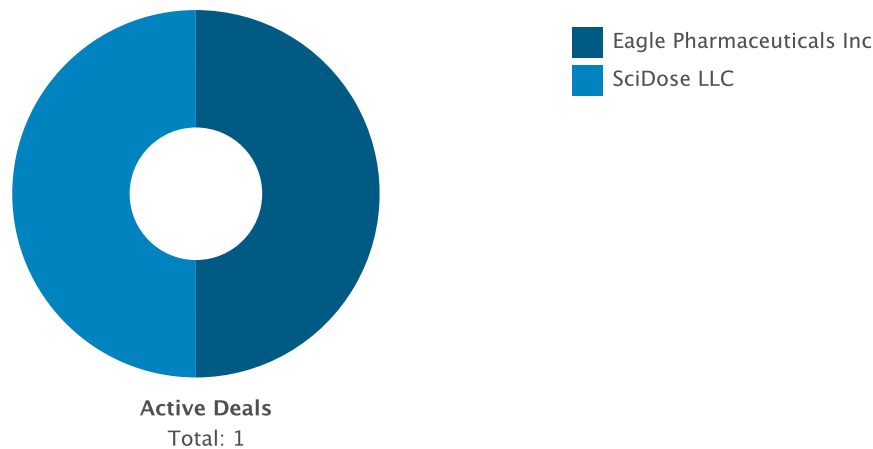
| Names | Type |
|---|------|
| undisclosed compounds (cancer), SciDose/Eagle Pharmaceuticals | |

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DEALS

Deals by Parent Company Chart

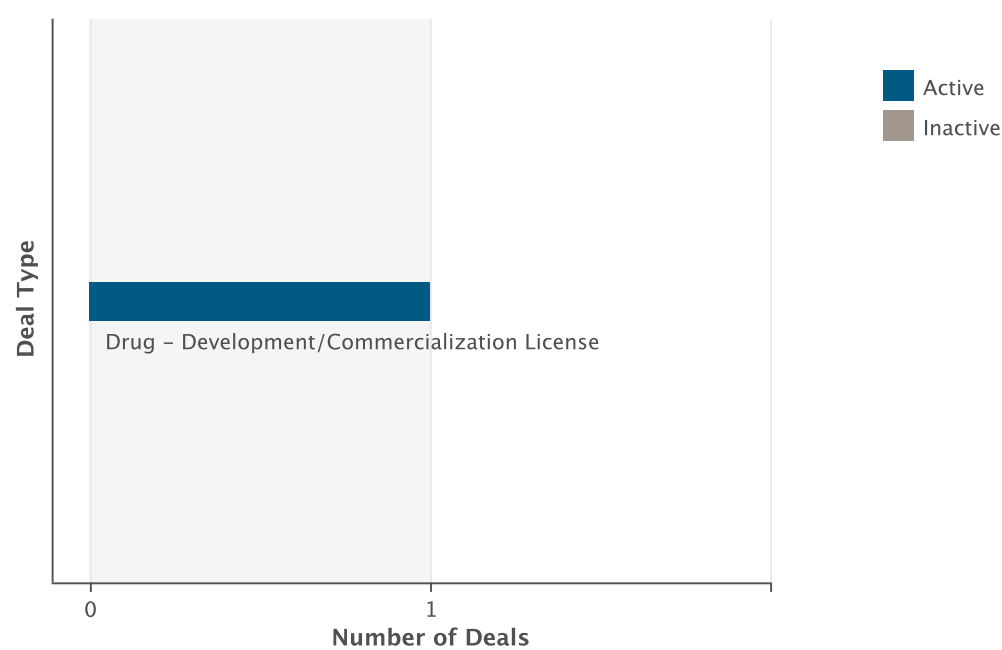


Deals by Parent Company Table

| Company Name | Principal | | Partner | | Total |
|---------------------------|-----------|----------|---------|----------|-------|
| | Active | Inactive | Active | Inactive | |
| Eagle Pharmaceuticals Inc | 0 | 0 | 1 | 0 | 1 |
| SciDose LLC | 1 | 0 | 0 | 0 | 1 |

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Deals by Type Chart



Deals by Type Table

| Deal Type | Active | Inactive | Total |
|--|--------|----------|-------|
| Drug - Development/Commercialization License | 1 | 0 | 1 |

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ready-to-dilute pemetrexed (iv, non-small cell lung cancer/mesothelioma), Eagle Pharmaceuticals

ready-to-dilute pemetrexed (iv, non-small cell lung cancer/mesothelioma), Eagle Pharmaceuticals SNAPSHOT

| | |
|-----------------------------|---|
| Drug Name | ready-to-dilute pemetrexed (iv, non-small cell lung cancer/mesothelioma), Eagle Pharmaceuticals |
| Key Synonyms | pemetrexed |
| Originator Company | Eagle Pharmaceuticals Inc |
| Active Companies | Eagle Pharmaceuticals Inc |
| Inactive Companies | |
| Highest Status | Discovery |
| Active Indications | Metastatic non small cell lung cancer;Mesothelioma |
| Target-based Actions | DHFR inhibitor;Thymidylate synthase inhibitor;Transferase inhibitor |
| Other Actions | Anticancer antimetabolite |
| Technologies | Small molecule therapeutic;Intravenous formulation;Infusion |
| Last Change Date | 09-Apr-2014 |

ready-to-dilute pemetrexed (iv, non-small cell lung cancer/mesothelioma), Eagle Pharmaceuticals DEVELOPMENT PROFILE

SUMMARY

Eagle Pharmaceuticals is investigating EP-5101, a ready-to-dilute (RTD) formulation of pemetrexed, an antifolate antineoplastic agent that exerts its action by disrupting folate-dependent metabolic processes essential for cell replication, for the potential treatment of locally advanced or metastatic non-small cell lung cancer and mesothelioma. By December 2013, formulation studies had been completed. In March 2014, the program was listed as being in preclinical formulation and toxicology studies.

ready-to-dilute pemetrexed (iv, non-small cell lung cancer/mesothelioma), Eagle Pharmaceuticals DEVELOPMENT STATUS

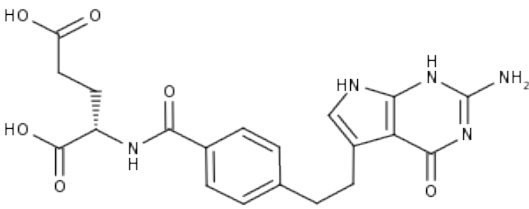
CURRENT DEVELOPMENT STATUS

| Company | Indication | Country | Development Status | Date |
|---------------------------|---------------------------------------|---------|--------------------|-------------|
| Eagle Pharmaceuticals Inc | Mesothelioma | US | Discovery | 20-Dec-2013 |
| Eagle Pharmaceuticals Inc | Metastatic non small cell lung cancer | US | Discovery | 20-Dec-2013 |

ready-to-dilute pemetrexed (iv, non-small cell lung cancer/mesothelioma), Eagle Pharmaceuticals CHEMICAL STRUCTURES

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| | |
|--|-------------------|
| CAS Registry Number: | Confidence Level: |
| 137281-23-3 | 1 |
|  | |
| Name | Type |
| pemetrexed | INN |
| EP-5101 | Research Code |

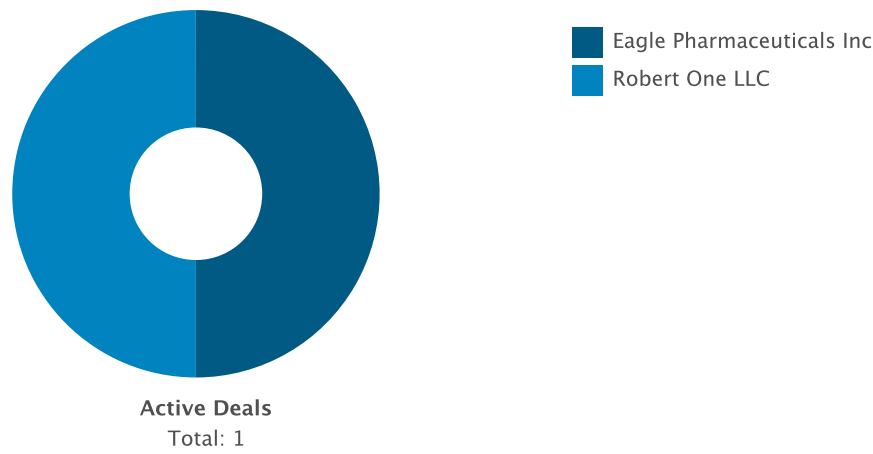
ready-to-dilute pemetrexed (iv, non-small cell lung cancer/mesothelioma), Eagle Pharmaceuticals
DRUG NAMES

| | |
|---|---------------|
| Names | Type |
| EP-5101 | Research Code |
| pemetrexed | INN |
| ready-to-dilute pemetrexed (iv, non-small cell lung cancer/mesothelioma), Eagle Pharmaceuticals | |

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DEALS

Deals by Parent Company Chart



Deals by Parent Company Table

| Company Name | Principal | | Partner | | Total |
|---------------------------|-----------|----------|---------|----------|-------|
| | Active | Inactive | Active | Inactive | |
| Eagle Pharmaceuticals Inc | 0 | 0 | 1 | 0 | 1 |
| Robert One LLC | 1 | 0 | 0 | 0 | 1 |

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Deals by Type Chart



Deals by Type Table

| Deal Type | Active | Inactive | Total |
|---------------------------|--------|----------|-------|
| Patent - Exclusive Rights | 1 | 0 | 1 |

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