

Otonomy Inc

CORTELLIS COMPANY DETAILED PIPELINE REPORT

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

Publication Date: 15-Dec-2014

THOMSON REUTERS

3 Times Square New York, New York 10036 United States

Tel: +1 646 223 4000

thomsonreuters.com



ABOUT CORTELLIS COMPANY DETAILED PIPELINE REPORT

Thomson Reuters provides the knowledge, tools, and expertise to help support drug discovery and development activities, IP portfolio optimization, identification of licensing and partnering opportunities, delivery of successful regulatory submissions, and the ability to keep current with the rapidly-changing pharmaceutical and chemical markets, supporting informed, early decisions.

This report was created by Thomson Reuters, using information from *Thomson Reuters Cortellis*™ for *Competitive Intelligence*; a comprehensive, proven intelligence solution that leverages the most accurate, complete, and widely respected drug pipeline information. From drug discovery and development activities to patent reports, the latest deals, and partnering opportunities, *Cortellis* can provide the confidence to make the most informed business decisions, faster. *Cortellis for Competitive Intelligence* provides accurate and validated information on pharmaceutical and biotechnology companies globally, their drug pipelines, deals, patents, and clinical trials, plus breaking industry news and conference coverage. All contained in one simple, highly intuitive research platform.

Cortellis Company Detailed Pipeline reports are the second in a series of that track pharmaceutical and biotechnology companies worldwide. All Cortellis for Competitive Intelligence content is subject to the most comprehensive editorial review process available, conducted by scientists, pharma professionals, regulatory experts, and generics specialists. Featuring timely drug pipeline information expertly uncovered and integrated from over 400 global meetings each year, you'll always be on top of the latest developments.

Chosen by leading life sciences companies, their executives and investors, *Cortellis for Competitive Intelligence* accelerates your deal-making and gives you timely insights on the development landscape.

Discover undiscovered opportunities in drug development and licensing faster with *Thomson Reuters Cortellis™ for Competitive Intelligence*

DISCLAIMER

The information contained in this report is based on sources believed to be correct but Thomson Reuters does not guarantee the accuracy, timeliness, or completeness of this information. Opinions, if any, are those held by the author of any individual report or article at the time of initial publication and do not necessarily reflect the views of Thomson Reuters.

Information in this report on companies is intended for reference use only, and does not constitute a recommendation to buy or sell any particular security or other investment and does not constitute an offer to buy from or sell to any particular investor. Any company or securities mentioned in this report may not be suitable for any particular investor, depending on that investor's financial position and needs.



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 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.

THOMSON REUTERS

TABLE OF CONTENTS

Company Overview	5
Company Profile	6
Product Portfolio Summary	6
Product Portfolio Drug Pipeline Detail	S
Phase 3 Clinical	10
Phase 2 Clinical	12
Discovery	17



Otonomy Inc

COMPANY OVERVIEW

O	
Company Name	Otonomy Inc
Parent Company Name	Otonomy Inc
Website	http://www.otonomy.com/
Country	US
Number of Drugs in Active Development	3
Number of Inactive Drugs	0
Number of Patents as Owner	12
Number of Patents as Third Party	0
Number of Deals	2
Key Indications	Tinnitus, Ear disease, Meniere disease, Otitis media, Hearing disorder, Chronic sinusitis, Fungal infection, Hearing loss, Nasal polyps, Dizziness, Epistaxis, Inner ear disease, Nasopharyngeal carcinoma, Nose disease, Otitis, Presbycusis, Sensorineural hearing loss, Vertigo
Key Target-based Actions	Glucocorticoid agonist,DNA gyrase inhibitor,Topoisomerase IV inhibitor,NMDA receptor antagonist,Metabotropic glutamate receptor 1 modulator,Metabotropic glutamate receptor 5 modulator,Metabotropic glutamate receptor modulator
Key Technologies	Otic gel formulation, Small molecule therapeutic, Sustained release formulation, Otic formulation, Steroid, Injectable controlled release formulation, Antibiotic, Injectable formulation, Parenteral formulation unspecified, Drug combination, Formulation preservation, Liquid-based formulation technology, Nanoparticle formulation

COMPANY PROFILE

SUMMARY

Otonomy, founded in 2008, is an emerging biopharmaceutical company dedicated to developing and marketing therapeutics designed specifically for the middle and inner ear.

FINANCIAL

In October 2014, Otonomy was added to the Russell 2000 Index.

In July 2014, the company filed a registration statement on Form S-1 for the proposed initial public offering. In August 2014, the company priced its initial public offering of 6,250,000 shares of common stock, at \$16 each. The underwriters were granted a 30-day option to purchase up to an additional 937,500 shares of common stock. At that time, the shares began trading on NASDAQ global select market under the symbol "OTIC". In September 2014, the underwriters purchased the additional 937,500 shares at \$16 each. The company raised \$104 million through the offering.

In April 2014, the company raised \$49 million from an oversubscribed series D financing round.

In September 2013, Otonomy raised \$45.9 million in a series C financing round.

In August 2010, the company raised \$38.5 million from a series B financing round.

In June 2010, the company raised \$10 million from a series A financing round.

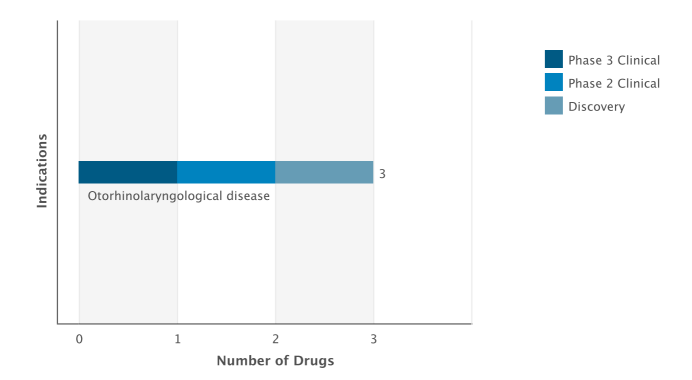
THOMSON REUTERS

PRODUCT PORTFOLIO SUMMARY

DRUGS

Drugs by Indication

Active Drugs by Indication Chart



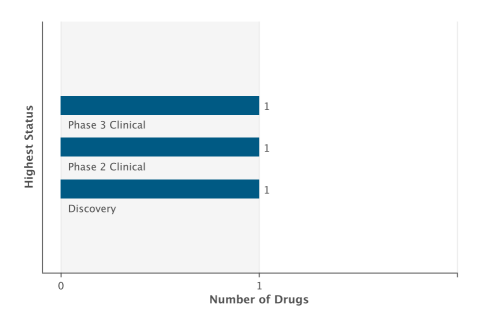
Drugs by Indication Table

Indication	Active	Inactive	Total
Otorhinolaryngological disease	3	0	3

THOMSON REUTERS

Drugs by Highest Status

Active Drugs by Highest Status Chart



Drugs by Highest Status Table

Development Status	Number of Drugs
Phase 3 Clinical	1
Phase 2 Clinical	1
Discovery	1

DEALS

Deal Type	Principal		Partner		Total
	Active	Inactive	Active	Inactive	
Drug - Asset Divestment	0	0	1	0	1
Drug - Early Research/Development	0	0	1	0	1

CLINICAL TRIALS

Trials by Condition Studied

Condition Studied	Ongoing	All
Otorhinolaryngological disease	1	6



Trials by Phase

Phase	Ongoing	All
Phase 3	0	2
Phase 2	1	2
Phase 1	0	2

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	1	0	1
Immune disorder	1	0	1
Neoplasm	1	0	1
Mouth disease	1	0	1
Neurological disease	4	0	4
Respiratory disease	1	0	1
Infectious disease	1	0	1
Injury	1	0	1
Inflammatory disease	1	0	1
Otorhinolaryngological disease	8	0	8

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

THOMSON REUTERS

PRODUCT PORTFOLIO DRUG PIPELINE DETAIL

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

ciprofloxacin (intratympanic sustained release gel, otitis media), Otonomy

ciprofloxacin (intratympanic sustained release gel, otitis media), Otonomy SNAPSHOT

Drug Name	ciprofloxacin (intratympanic sustained release gel, otitis media), Otonomy
Key Synonyms	ciprofloxacin;AuriPro
Originator Company	Otonomy Inc
Active Companies	Otonomy Inc
Inactive Companies	
Highest Status	Phase 3 Clinical
Active Indications	Otitis media
Target-based Actions	DNA gyrase inhibitor;Topoisomerase IV inhibitor
Other Actions	Antibacterial;Bacterial nucleic acid synthesis inhibitor
Technologies	Sustained release formulation;Injectable formulation;Antibiotic;Small molecule therapeutic;Parenteral formulation unspecified;Otic gel formulation
Last Change Date	22-Oct-2014

ciprofloxacin (intratympanic sustained release gel, otitis media), Otonomy DEVELOPMENT PROFILE

SUMMARY

Otonomy is developing AuriPro (previously OTO-201; OTO-203), a sustained release gel formulation of ciprofloxacin, delivered using intratympanic injection, for the potential treatment of otitis media,,. In November 2013, two phase III trial were initiated in the US and Canada. In December 2013, dosing of the first patient was initiated,; In May 2014, enrollment was completed. In July 2014, positive data were reported. In October 2014, the company was planning to file for an NDA in the first half of 2015. In November 2013, the company was seeking to outlicense the drug.

ciprofloxacin (intratympanic sustained release gel, otitis media), Otonomy DEVELOPMENT STATUS

CURRENT DEVELOPMENT STATUS

Company	Indication	Country	Development Status	Date
Otonomy Inc	Otitis media	Canada	Phase 3 Clinical	11-Dec-2013
Otonomy Inc	Otitis media	US	Phase 3 Clinical	11-Nov-2013

ciprofloxacin (intratympanic sustained release gel, otitis media), Otonomy CHEMICAL STRUCTURES



CAS Registry Number:	Confidence Level:
85721-33-1	1
t HN N	N O OH
Name	Туре
ciprofloxacin	BANN; INN; USAN
Pulmaquin	Trade Name
OBM-A-03	Research Code
OTO-201	Research Code
Cipro NDS	

ciprofloxacin (intratympanic sustained release gel, otitis media), Otonomy DRUG NAMES

Names	Туре
ciprofloxacin	USAN, BANN, INN
ciprofloxacin (intratympanic sustained release gel, otitis media), Otonomy	
AuriPro	Trade Name
OTO-201	Research Code
OTO-203	Research Code

ciprofloxacin (intratympanic sustained release gel, otitis media), Otonomy CLINICAL TRIALS

Trials by Phase and Condition Studied

	se 4 nical		se 3 nical		se 2 iical		se 1 nical	Pha Unspe	ase ecified	То	tal
On- going	All	On- going	All								
Otitis media											
0	0	0	1	0	0	0	1	0	0	0	2



Total Trials by Phase and Status

	se 4 iical	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	1	0	0	0	1	0	0	0	2

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



dexamethasone (sustained release/intratympanic injection, ear disorders), Otonomy

dexamethasone (sustained release/intratympanic injection, ear disorders), Otonomy SNAPSHOT

Drug Name	dexamethasone (sustained release/intratympanic injection, ear disorders), Otonomy
Key Synonyms	dexamethasone
Originator Company	Otonomy Inc
Active Companies	Otonomy Inc
Inactive Companies	
Highest Status	Phase 2 Clinical
Active Indications	Meniere disease;Ear disease
Target-based Actions	Glucocorticoid agonist
Other Actions	Steroidal anti-inflammatory
Technologies	Injectable controlled release formulation;Otic formulation;Small molecule therapeutic;Otic gel formulation;Steroid
Last Change Date	10-Oct-2014

dexamethasone (sustained release/intratympanic injection, ear disorders), Otonomy DEVELOPMENT PROFILE

SUMMARY

Otonomy is developing OTO-104, a local-acting, sustained-release thermosensitive gel formulation of dexamethasone, delivered via intratympanic injection, for the potential injectable treatment of ear diseases, including Meniere disease. In November 2013, a phase II/III trial for Meniere disease was ongoing. In February 2011, the company planned to initiate clinical trials for noise- and chemotherapy-induced hearing loss. In August 2013, the company was advancing the formulation into late-stage clinical trials. In October 2014, a multiple dose safety study of OTO-104 was initiated in UK and at that time, the company planned to start clinical development activities outside of North America. In September 2009, the company was seeking to outlicense the drug outside the US; in November 2013, this was still the case.

dexamethasone (sustained release/intratympanic injection, ear disorders), Otonomy DEVELOPMENT STATUS

CURRENT DEVELOPMENT STATUS

Company	Indication	Country	Development Status	Date
Otonomy Inc	Meniere disease	Canada	Phase 2 Clinical	11-Dec-2013
Otonomy Inc	Meniere disease	US	Phase 2 Clinical	11-Sep-2011
Otonomy Inc	Meniere disease	UK	Clinical	09-Oct-2014
Otonomy Inc	Ear disease	US	Discovery	21-Sep-2009



dexamethasone (sustained release/intratympanic injection, ear disorders), Otonomy CHEMICAL STRUCTURES

CAS Registry Number:	Confidence Level:
50-02-2	1
	0
но▲	V JumoH OH
	H >
	H
0	J "
Name	Туре
dexamethasone	BANN; INN
Ozurdex	Trade Name
Surodex	Trade Name
CD-102	Research Code
TLC-x99	Research Code
120 000	research odde
OTX-DP	Research Code
IBI-10090	Research Code
101-10090	Research Code
EGP-437	Research Code
OTO-104	Research Code
010-104	Nescalul Coue
ISV-305	Research Code
Opposit	
Oncocort	

dexamethasone (sustained release/intratympanic injection, ear disorders), Otonomy DRUG NAMES

Names	Туре
OTO-104	Research Code
dexamethasone	BANN, INN
dexamethasone (sustained release/intratympanic injection, ear disorders), Otonomy	

dexamethasone (sustained release/intratympanic injection, ear disorders), Otonomy CLINICAL TRIALS



Trials by Phase and Condition Studied

	se 4 lical		se 3 nical		se 2 nical		se 1 nical	Pha Unspe		То	tal
On- going	All	On- going	All	On- going	All	On- going	All	On- going	All	On- going	All
Meniere o	Meniere disease										
0	0	0	0	1	2	0	1	0	0	1	3

Total Trials by Phase and Status

	Phase 4 Phase 3 Clinical 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	1	2	0	1	0	0	1	3

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 Clinica

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

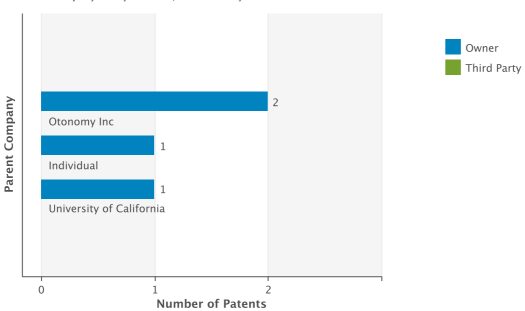
THOMSON REUTERS

dexamethasone (sustained release/intratympanic injection, ear disorders), Otonomy DEALS AND PATENTS

PATENTS

Patents by Parent Company Chart

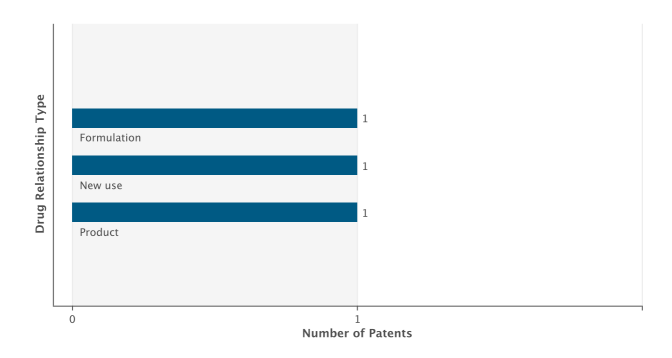
Chart displayed by Owner/Third Party



Patents by Parent Company Table

Company Name	As Owner	As Third Party	Total
Otonomy Inc	2	0	2
Individual	1	0	1
University of California	1	0	1

Patents by Drug Relationship Type Chart



Patents by Drug Relationship Type Table

Drug Relationship	Total
Formulation	1
New use	1
Product	1

OTO-311

OTO-311 SNAPSHOT

Drug Name	OTO-311
Key Synonyms	gacyclidine
Originator Company	DURECT Corp
Active Companies	Otonomy Inc
Inactive Companies	NeuroSystec Corp;DURECT Corp
Highest Status	Discovery
Active Indications	Tinnitus
Target-based Actions	NMDA receptor antagonist
Other Actions	Analgesic
Technologies	Injectable controlled release formulation;Drug implant;Sustained release formulation;Nanoparticle formulation injectable;Otic formulation;Small molecule therapeutic;Local formulation unspecified
Last Change Date	06-Nov-2014

OTO-311 DEVELOPMENT PROFILE

SUMMARY

Otonomy under license from NeuroSystec, (under exclusive license from DURECT), is developing OTO-311 (NST-001), a squalene-based nanoparticle formulation of gacyclidine, an NMDA receptor antagonist delivered directly into the cochlear using Otonomy's proprietary drug delivery technology (previously delivered using Neuroject implantable continuous delivery system), for the potential injectable treatment of tinnitus,. In November 2013, the drug was listed as being in preclinical development and at that time, the company was seeking to outlicense the drug. In November 2014, Otonomy was planning clinical trials for the following year.

OTO-311 DEVELOPMENT STATUS

CURRENT DEVELOPMENT STATUS

Company	Indication	Country	Development Status	Date
Otonomy Inc	Tinnitus	US	Discovery	06-Nov-2013
DURECT Corp	Tinnitus	France	Discontinued	21-Jun-2004
NeuroSystec Corp	Tinnitus	France	Discontinued	31-Oct-2013

OTO-311 CHEMICAL STRUCTURES



CAS Registry Number:	Confidence Level:			
68134-81-6	1			
	· s			
N.				
Name	Туре			
gacyclidine	INN			
GK-11	Research Code			
NST-001	Research Code			

OTO-311 DRUG NAMES

Names	Туре
NMDA antagonist (nanoparticle implant, tinnitus), NeuroSystec Corp	
NST-001	Research Code
gacyclidine	INN
gacyclidine (nanoparticle implant, tinnitus), NeuroSystec Corp	
NMDA receptor antagonist (nanoparticle implant, tinnitus), Otonomy	
NMDA receptor antagonist (nanoparticle implant, tinnitus), NeuroSystec Corp	
OTO-311	Research Code

OTO-311 CLINICAL TRIALS

Trials by Phase and Condition Studied

	se 4 nical		se 3 nical		se 2 nical		Phase 1 Clinical		ase ecified	То	tal
On- going	All	On- going	All	On- going	All	On- going	All	On- going	All	On- going	All
Tinnitus											
0	0	0	0	0	0	0	1	0	0	0	1



Total Trials by Phase and Status

	se 4 ical		se 3 lical		se 2 nical		Phase 1 Phase Clinical Unspecified		То	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

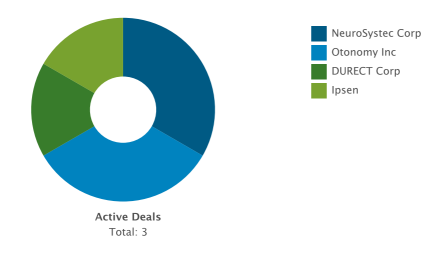
Includes Phase 2, Phase 2a, Phase 2b, Phase 1/2 (where enrolment count is 100 or over), Phase 2/3 (where enrolment count is under 300 or not specified)

Phase 1 Clinical

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

OTO-311 DEALS AND PATENTS

DEALS Deals by Parent Company Chart

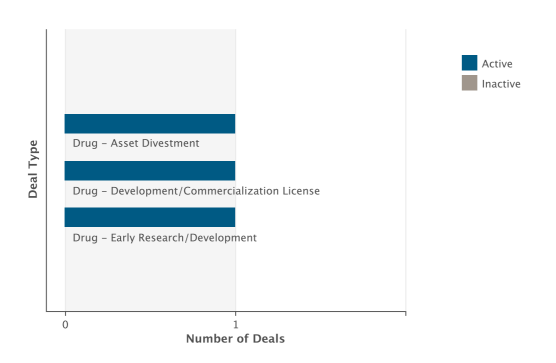




Deals by Parent Company Table

Company Name	Principal Active Inactive		Partner Active Inactive		Total
Otonomy Inc	0	0	2	0	2
NeuroSystec Corp	1	0	1	0	2
Ipsen	1	0	0	0	1
DURECT Corp	1	0	0	0	1

Deals by Type Chart



Deals by Type Table

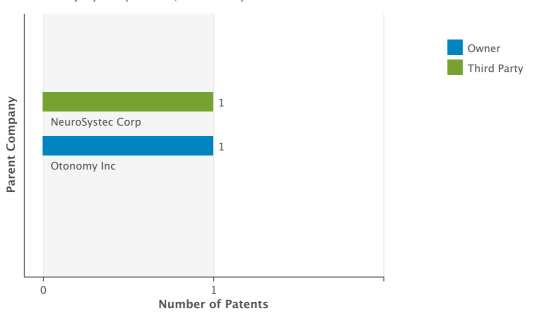
Deal Type	Active	Inactive	Total
Drug - Early Research/Development	1	0	1
Drug - Asset Divestment	1	0	1
Drug - Development/Commercialization License	1	0	1



PATENTS

Patents by Parent Company Chart

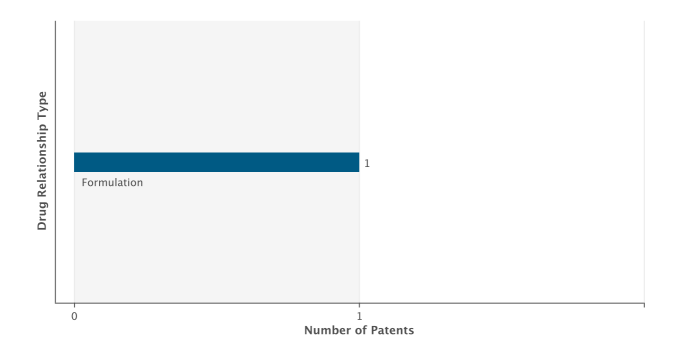
Chart displayed by Owner/Third Party



Patents by Parent Company Table

Company Name	As Owner	As Third Party	Total
Otonomy Inc	1	0	1
NeuroSystec Corp	0	1	1

Patents by Drug Relationship Type Chart



Patents by Drug Relationship Type Table

Drug Relationship	Total
Formulation	1

This report was created by Thomson Reuters, using information from *Thomson Reuters Cortellis*™ *for Competitive Intelligence*; a comprehensive, proven intelligence solution that leverages the most accurate, complete, and widely respected drug pipeline information.

For more information about *Cortellis for Competitive Intelligence*, visit: http://cortellis.thomsonreuters.com/cortellis_for_you/?cid=thomsonone.

For subscription information, e-mail scientific.lifesciences@thomsonreuters.com.

© 2012 Thomson Reuters. All rights reserved. Republication or redistribution of Thomson Reuters content, including by framing or similar means, is prohibited without the prior written consent of Thomson Reuters. 'Thomson Reuters' and the Thomson Reuters logo are registered trademarks and trademarks of Thomson Reuters and its affiliated companies.

THOMSON REUTERS