Confirming Ipsen's specialist care globalisation

28^{HP} Annual JP Morgan Healthcare Conference San Francisco, January 11-13, 2010 Mr. Stéphane Thiroloix - EVP Corporate Development Dr. Jacques-Pierre Moreau – Chief Scientific Advisor Mr. Pierre Kemula – Investor Relations Manager





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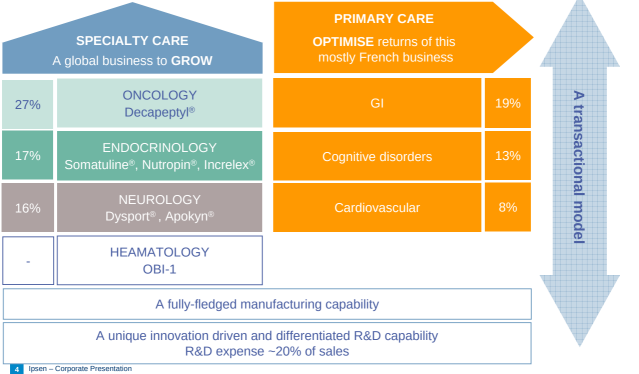
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Introduction



SIPSEN Innovation for patient care

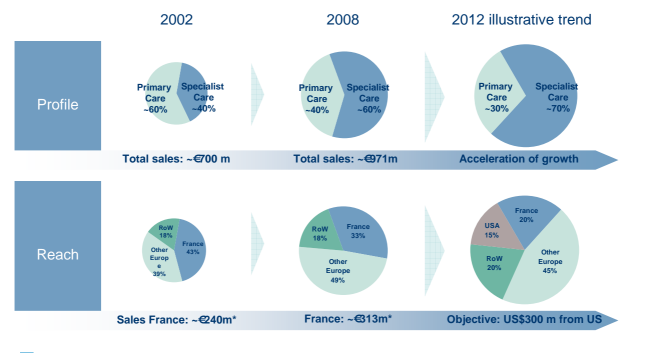
Ipsen today : a global, innovation driven, specialty pharma



* % are calculated on 2008 total Group Drug Sales of $\,{\in}936$ million

A reinforced profile and reach

SIPSEN ///////



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* Excludes sales of Ginkor Fort (€61 million in 2002, €14 million in 2008) specialist care and primary care relative weights are expressed as a % of total Drug sales



An increasingly transactional model





Ipsen today....

- Resilience of business in a difficult macro-economic environment
- A strong and profitable specialty care growth engine
- Substantial growth opportunities through globalization and US entry
- A rich and well balanced R&D pipeline, with potential blockbusters
- A strong cash flow generation and balance sheet

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Truly Differentiated R&D Capabilities





Defining Ipsen's competitive edge in R&D

Hormones provide well defined templates with matching targets both novel or validated

Resident know how based on the integration of basic discovery technologies

Technologies	Medicinal chemistry	Delivery systems
Target identification, validation and drugability based on clinical observations supported byomics technologies	Steroids peptides, proteins engineering aiming at enhanced efficacy, potency, selectivity and safety over the endogenous hormone	Emphasis on improved pharmacological properties, optimization of dosing regimen and improved patients compliance and convenience

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Example 1 : Somatuline[®] Depot, an improved presentation

	Sandostatin LAR®	Somatuline [®] Autogel [®]			
Administration	2.0 ml Intramuscular	0.3 ml – 0.5 ml Subcutaneous			
Presentation	Powder vial + solvent filled syringe + 2 needles	Pre-filled syringe			
Injection technique	10 steps needed to reconstitute	Ready to use Self administration*			



For what reasons would you prescribe Somatuline® Depot to your acromegaly patients?**

Pre-filled sy	ringe / n	o recons	titution n	eeded				879
More conver	nient be	cause the	e patient	can self i	nject		83	%
Saves staff t	ime and	resource	es (self-inje	ction possible	e at home)	65%		
Improved pa (less injection site				ller volume)	61%			
			%	of physicial	าร			

* In selected countries

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** Study Sample: A total of 50 US endocrinologists completed a 30-minute online questionnaire between April 4 - 17, 2008 25 High Volume Endocrinologists: Endocrinologists who see 11 or more acromegaly patients in a year 25 Low Volume Endocrinologists: Endocrinologists who see between 5-10 acromegaly patients in a year



Example 2 : a unique technology convergence, taspoglutide

Once-a-week injection

- Equal / greater potency compared to native compound
- Extended metabolic half-life, 22x more stable in plasma
- Complete retention of incretin properties
- Strong patent positions



Expected needle gauge

- (LAR) → 23G
 Quarter inch long
- Taspoglutide Liquid SRF → 29G
 Insulin type needle for subcutaneous injection

50 to 300 µl of highly concentrated aqueous solution devoid of excipient

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Taspoglutide Toxicology Roche/Ipsen perspective on GLP-1s and C-cell thyroid tumors

Results	 Results of carcinogenicity studies support the ongoing clinical development of taspoglutide; These results apply to mice as well
Disclosure	 We will share the results of the 2 year toxicology studies with the regulatory authorities first and may share the data with the public at a later point in time

Roche Reminder - ADA 2009 last toxicology update Roche/Ipsen perspective on GLP-1s and C-cell thyroid tumors

Preliminary incidence	 Taspoglutide carcinogenicity program ongoing Preliminary incidence of C-cell hyperplasia and adenoma was observed in a small number of animals Safety data not final yet; aiming for high-quality analysis to obtain reliable results
Disclosure	 Clinical relevance of proliferative lesions in rodents under investigation Working hypothesis: C-cell effects are rodent-specific Appropriate pre-clinical mechanistic studies underway Calcitonin monitoring in phase III

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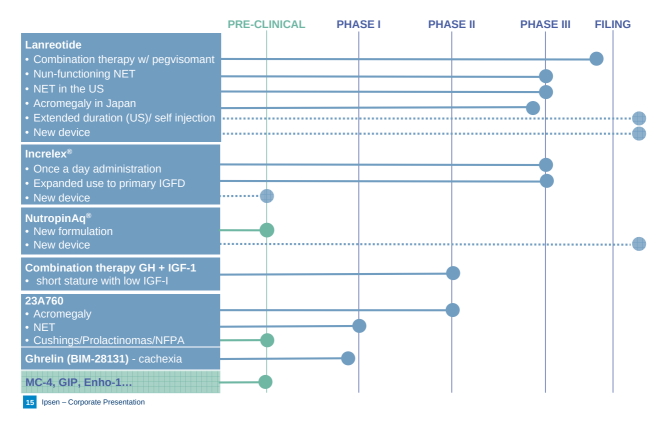
A rich and promising pipeline



•emerge



A rich endocrinology pipeline





A promising Oncology pipeline

	PRE CL	INICAL	PHA	SE I PI	HASE II	PHAS	SE III	FILING
Decapeptyl®								
• 6 M formulation								
Toremifene citrate 80 mg Treatment ADT induced side effects 								
• 20 mg HG PIN *						_		
BN-80915 (Diflomotecan)								
Advanced Metastatic Cancers								
BN-80927 (Elomotecan)								
Advanced Metastatic Cancers								
BN-83495 Advanced Breast & Prostate Cancer 								
Gynecological Cancers								
STX-140 (Angiomates)								
BIM-46187 (G-protein inhibitor)								
IRC-08364 (CDC 25 inhibitor)	•• Li	censed o	ut to L	Debiopharm				
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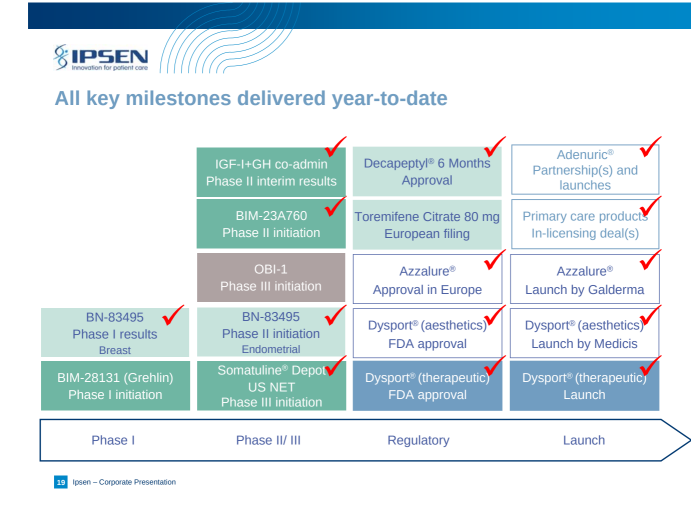






A track record for consistently delivering on strategic milestones

LCM	Decapeptyl [®] - 3 months formulation - Launches in the UK and Germany ~ 48% of European market - Debiopharm agreement for 6M approval
	Taspoglutide Ph I Ph III results Ph III
Pipeline Progress	OBI-1 — Ph I — Octagen Assets _ Ph III - Ph III
3	Adenuric [®] — Ph III — Filing – Approval — partnership →
Optimization	Ginkor [®] Fort divestment Adrovance [®] Exforge [®] co-marketing co-promotion
US entry	Strategic intent - First step of US entry (endocrinology) - Somatuline® Depot approval Full fledge presence in the US (Endocrinology + Neurology) IGF-1 / GH - Tercica Acquisition - Ph II / III ->
	Toxin Aesthetics Medicis Filing Approval → Toxin Therapeutic Filing Approval →





2009 financial objectives confirmed

Drug sales growth of 7.0 to 9.0%

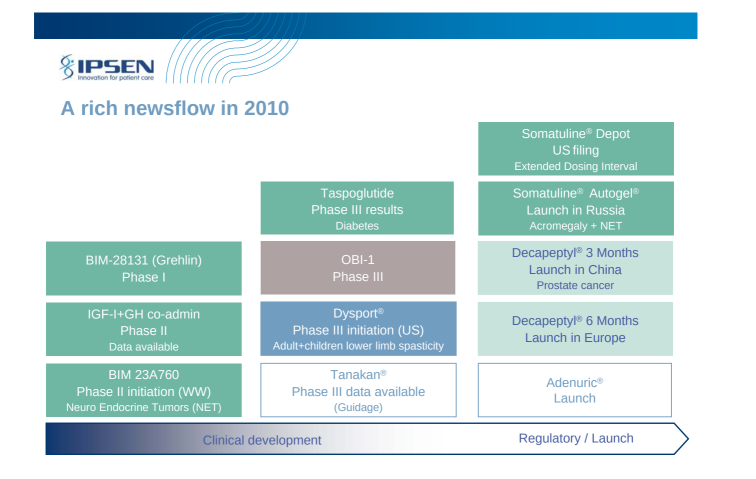
Other revenues¹ of approximately €80 million

Operating margin before goodwill allocation² of 17.0% - 17.5%

The above objectives include the full Kogenate® royalty stream

1- Defined as the total of milestone payments received under licence agreements, royalties received from third parties and other revenue (including for example co-promotion revenues)

> 2- These financial objectives do not include items resulting from purchase price accounting impacts related to the Group's transactions in North America

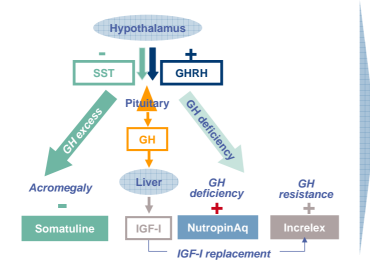


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An endocrinology franchise outgrowing competition



A unique focus on pituitary disorders and hormone dependent diseases



A strong franchise

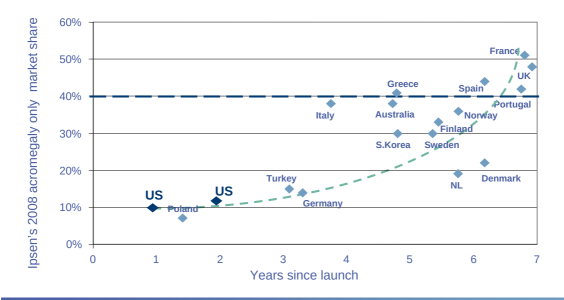
- A range of products addressing today Short Stature, Acromegaly and NET
 - High morbi-mortality
 - Debilitating pathologies
 - High unmet medical needs
- Somatuline[®], NutropinAq[®] and Increlex[®] contributed to ~16 % of 2008 Group sales, ie. ~ €158 million.
- A fast growing franchise: sales doubled in the past 3 years

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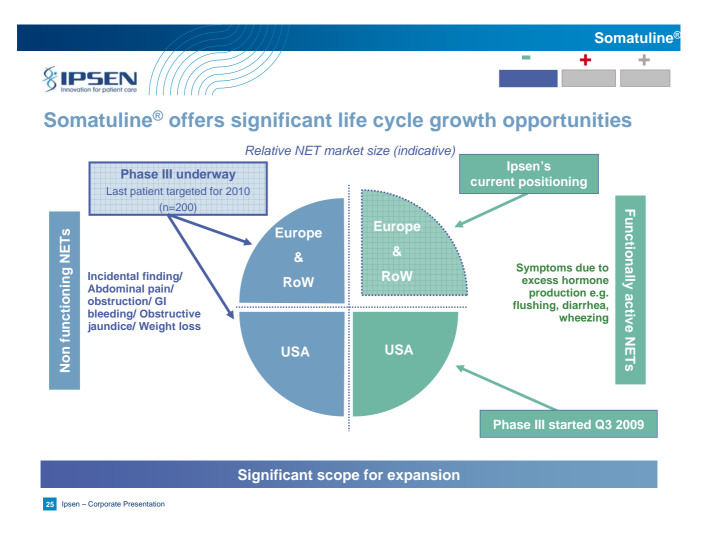
81PSEN //////



Somatuline[®] Depot is poised to grow and gain market share



Somatuline[®] market share is directly correlated to its time on market





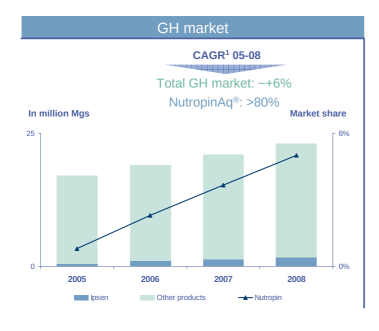
Increlex[®] in the US : steady performance with continued growth expectations

Physician demand	 Target audience : ~1,000 US paediatric endocrinologists Up to 20% of Rx come from new prescribers each month 2/3 of pediatric endocrinologists have prescribed Increlex[®]; 78% continued prescription
Reimbursement success	 ~ 90% of private and public covered lives have formulary access 75% Increlex patients approved upon final decision (similar to GH)
Patient experience	 Sharp increase in patients on Increlex[®] initially GH-naïve to 60% in '08 from 30% in '07 Dose increasing to appropriate targets, to 100 mcg/kg BID in '08 from 70 mcg/kg BID in '07 Younger patients initiated with Increlex[®], to average age at start of 10.0 years old in '08 from 11.5 years in '07

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NutropinAq[®] in Ipsen territories is steadily gaining market share



NutropinAq[®] attributes 1st liquid formulation launched WW A simple and user friendly pen An experienced post marketing

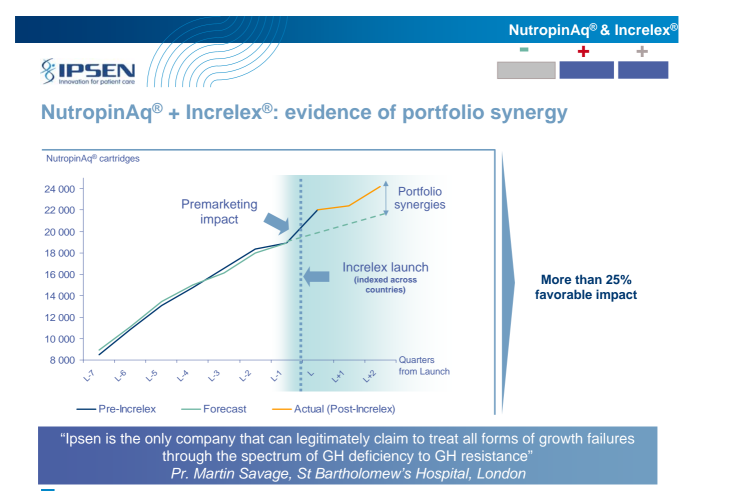
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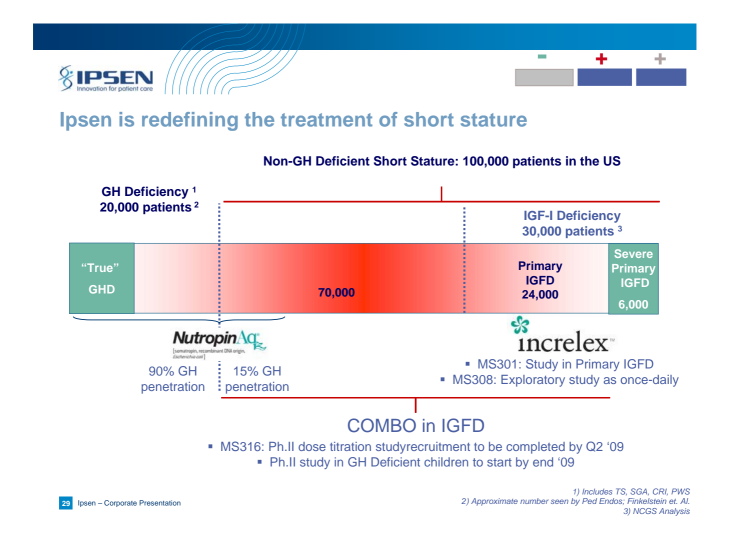
Source: Strategix

 A dedicated experienced and professional team

surveillance database

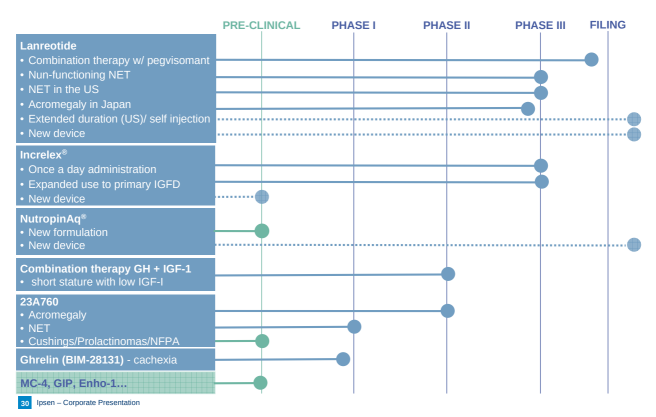
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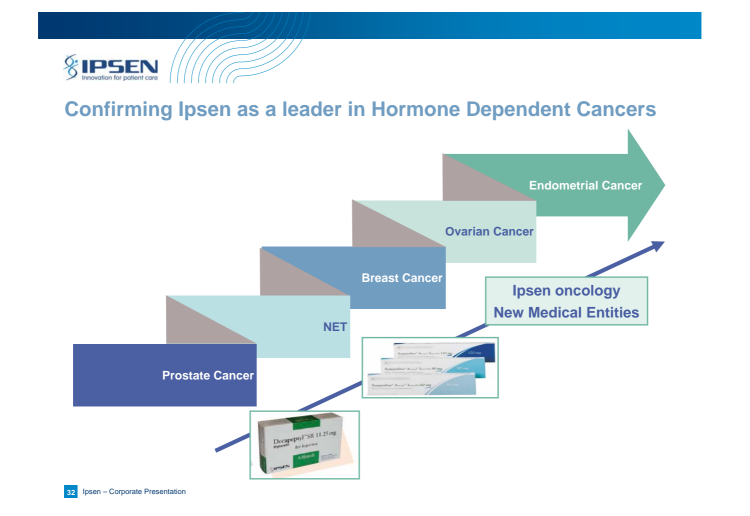


A rich endocrinology pipeline



Confirming Ipsen as a leader in the field of hormone dependent cancers





SIPSEN Innovation for patient care

Decapeptyl[®] 3 months formulation: a competitive product profile

Formulation and efficacy	 Marketed 1 month (1M) and 3 month (3M) formulations Maintenance of castrate testosterone levels at 3M in 98% of patients¹ At 3M, 91% decrease of PSA levels, showing tumor control 					
Local tolerance/ convenience	IM route of administration, good local toleranceInjection not visible for the patient					
Storage and reconstitution	 Stored at room temperature 5 steps reconstitution Safety needle system 					
	Competitor 1 Competitor 2 Competitor 3					
Formulation and efficacy	 Various formulations across territories : 1M formulation = 3,75mg or 7,5mg and 3M formulation = 11,25mg or 22,5mg Increased survival rate at 9 months in triptorelin group vs competitor 1² Conservation between 2 - 4° = needs to be warmed up before reconstitution Manual reconstitution to obtain SR Risk of nodules, abscess 					

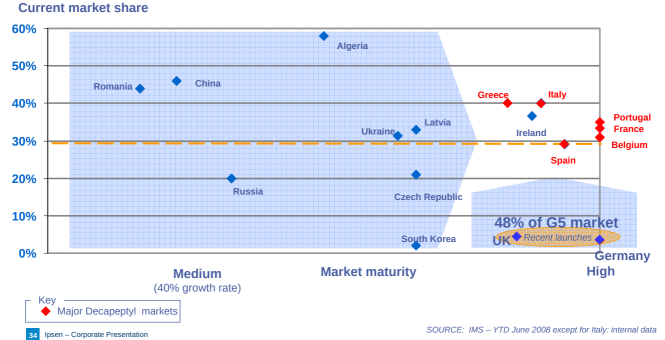
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SOURCE: French SmPC

REFERENCE . 1: Teillac, Horm Res, 2004, 252-58 2: Heyns, BJU Int, 2003, 226-231



Decapeptyl[®]: strong positions, and poised to grow



SIPSEN Decapeptyl[®] 6 month formulation: a more differentiated product profile

Efficacy	 Comparable efficacy to 1 and 3 months formulation Castration levels (testosterone) Disease control (PSA) 					
Local Tolerance	 Limited local side effects (6.7% of p 	patients)				
Storage and reconstitution	 Storage at room temperature (no n 5 Steps to reconstitute, change need 	eed to heat up before reconstitution) edle, and inject - IM route				
	o 1 (

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Reference 1: French SmPC ²Avis de la commission de transparence

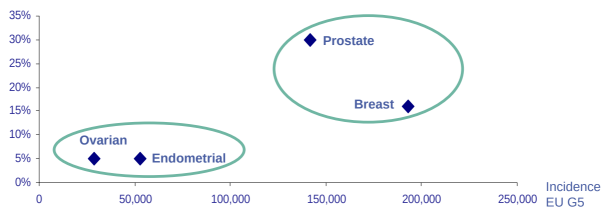


A promising Oncology pipeline

	PRE CI	INICAL	PHA	SE I F	PHASE II	PHAS	SE III	FILING
Decapeptyl®								
6 M formulation								T
Toremifene citrate 80 mg Treatment ADT induced side effects 								
• 20 mg HG PIN *								
BN-80915 (Diflomotecan)								
Advanced Metastatic Cancers								
BN-80927 (Elomotecan)								
Advanced Metastatic Cancers								
BN-83495								
Advanced Breast & Prostate CancerGynecological Cancers								
STX-140 (Angiomates)								
BIM-46187 (G-protein inhibitor)								
IRC-08364 (CDC 25 inhibitor)								
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SIPSEN Moving up to higher prevalence diseases and higher unmet medical needs





Ipsen New Medical Entities: multi targeted agents aiming at large markets as well as niche indications with large unmet medical needs BN-83495 is potentially a company transforming product

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SOURCE: deVita (2008), Datamonitor

Oristusane (BN-83495)



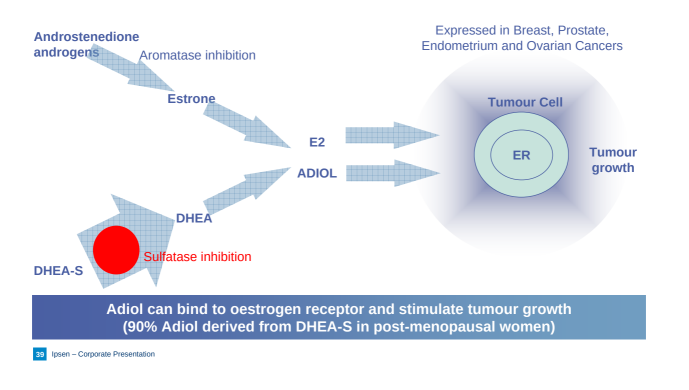
Oristusane : Moving forward in the development stages

Endometrial	 Phase II initiated : Post-menopausal women with advanced or recurrent endometrial cancer (80 patients) First patient dosed on November 25 					
Breast	 Phase I/ II on going in ER-positive metastatic breast cancer (35 patients) Optimal biological dose determined :40 mg once daily oral administration 95% inhibition of the target enzyme (STS) was achieved in peripheral blood mononuclear cells Additional 15 patients included to study target enzyme (STS) inhibition in cancerous cell 					
Prostate	 Phase II initiated – Dose escalation 					
Ovarian	 Course of action being defined 					
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Rationale for Sulfatase inhibitor development

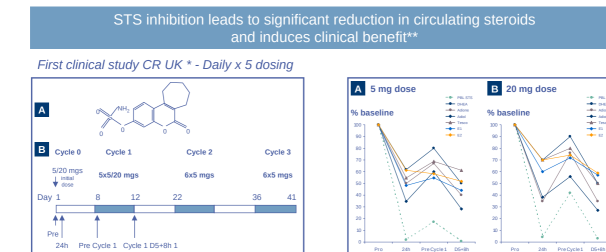
Inhibition of Androstenediol synthesis from DHEA-S



BN-83495



First clinical study in Breast Cancer patients



Pro

24h

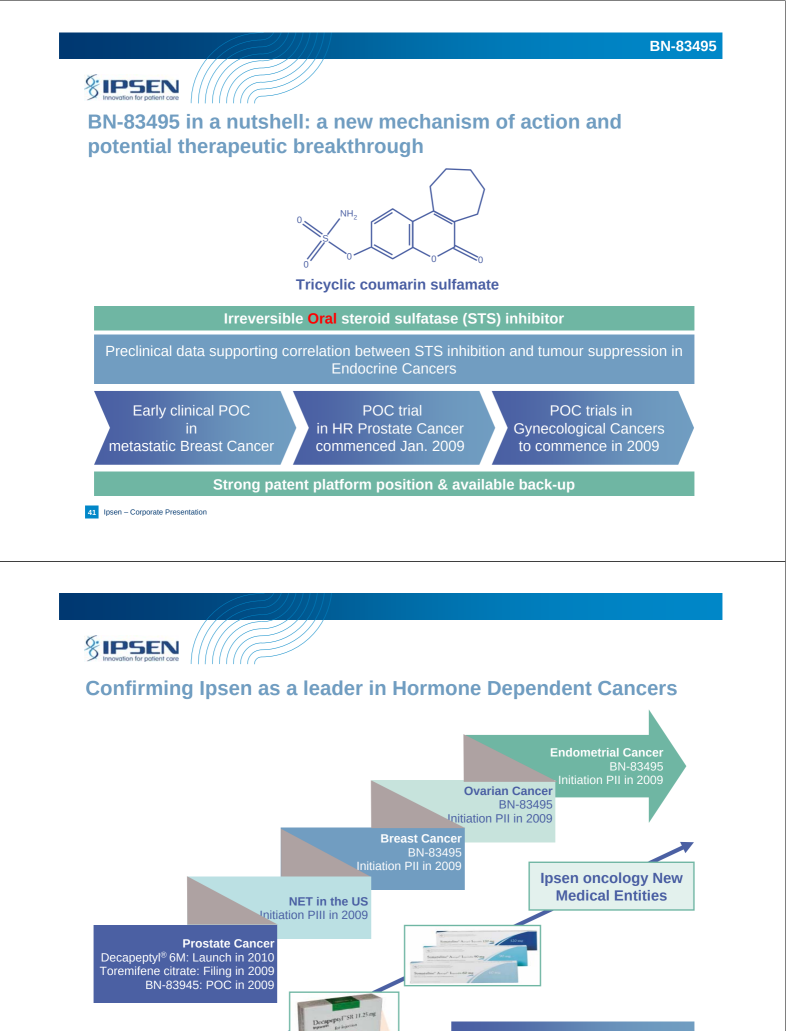
Pre Cycle 1 D5+8h

Next step: confirmation of the results in Metastatic Breast Cancer and exploration of the full range of hormonal dependent tumours

24h

Pre Cycle 1 D5+8h

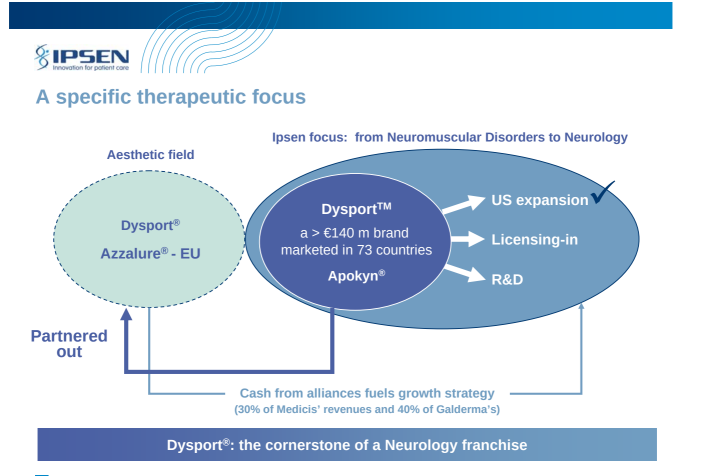
Pro

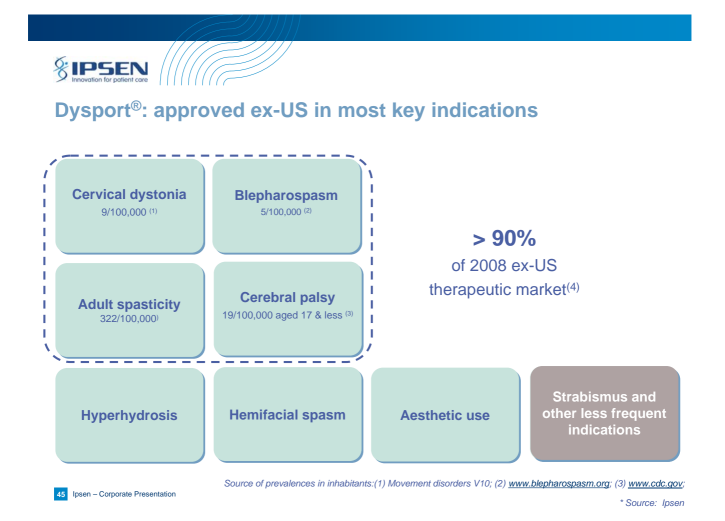


Near-term milestones...

From a Regional Neuromuscular Specialty to a Global Neurology Franchise

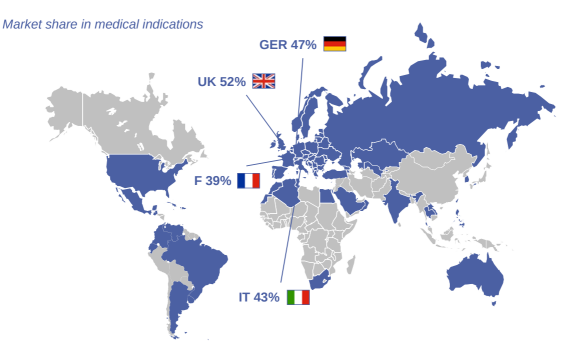


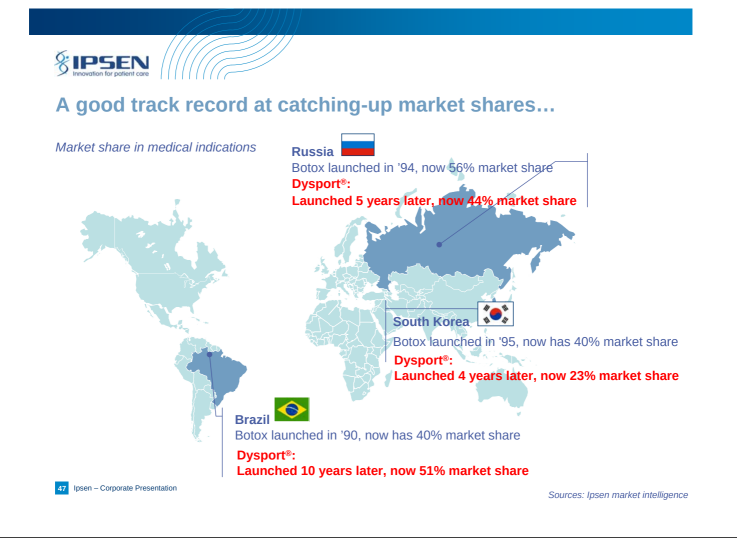






Dysport®: launched in 1991, approved in more than 75 countries







Dysport[®] in the US: a step further toward a global neurology franchise

- 1. Dysport®: a proven track record and field proven product
- 2. A true global product
- 3. A unique focus on medical use
- 4. Focus on US opportunity strong positioning with well prepared launch
 - Sound value proposition: the medical treatment alternative
 - Targeted and appropriate sales force
 - Managed care experience
- 5. Building up a neurology franchise leveraging the business development capability
- 6. Intense efforts in the discovery area

A focused haematology presence





An agent targeting both acquired and congenital hemophilia

Congenital hemophilia A with inhibitors to human FVIII

- Affects 1:4000 male births
- The development of neutralizing antibodies (inhibitors) to hFVIII following replacement therapy is a major complication
- Inhibitors develop in about 28% of severe patients and in between 3% to 13% of mild and moderate hemophilia A patients
- Patients no longer respond to hFVIII therapy

Acquired hemophilia Acquired factor VIII inhibitor

- Affects 1 to 2 individuals in 1,000,000, predominantly in older individuals
- A small proportion of younger patients may develop the disease, predominantly postpartum women
- Clinical manifestation is more severe and anatomically diverse than in congenital hemophilia A
- A mortality rate approaching 20%. Bleeding is often spontaneous or in response to minimal trauma

pFVIII is a promising treatment to stop bleeds in patients with inhibitors to hFVIII



Now preparing for phase 3...

2 prospective clinical trials, in liaison with Medical Community & Regulatory Agencies

Study in patients with acquired factor VIII inhibitor (acquired hemophilia)

Treatment of all acute bleeding episodes

Study in patients with congenital hemophilia A and inhibitors to hFVIII

Treatment of life or limb threatening bleeding episodes

Both will be of similar design Open label, non comparative prospective studies, with about 40 patients in each study

Standards setting: first ever prospective trial in acquired hemophilia

Protocols finalization and pre-phase 3 CMC consultations with regulatory agencies to be completed in H1 2009

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A highly specialized hospital product addressing unmet need

First biologics to conclude Phase 2 resulting from strategic biotechnology platform

Patent protection until 2023 in Europe and US

World-wide commercialization rights

Lean commercial infrastructure

A commercial potential in excess of US\$200 million

Fourth specialty therapeutic focus in Haematology

First half 2009 achievements





A strong commercial performance in the first half 2009

6.3% Drug sales growth, in line with our full-year objective

A solid 11.5% specialist care sales growth, with endocrinology up 32.7% year-on-year

Stabilisation in Eastern Europe, with Q2 sales up 1.0% year-on-year

Dynamic growth in the US, with Somatuline[®], Increlex[®] and Apokyn[®] generating \$23 million, up 33% Q2 over Q1



A strong profitability and cash generation

25.0% operating margin pre-goodwill allocation

A 'clean operating margin'* of 18.0%, compared with 21.6% a year ago

€147 m generated by operating activities, versus €124 m a year ago

€139 m net cash position as at June 30, 2009, post €203 m net cashed-out on US acquisitions in H2 08

NOTE: All margins expressed in % of sales

55 Ipsen – Corporate Presentation *Reported operating income excluding non-recurring elements (divestment of Ginkor Fort® & sale of land), purchase price accounting impacts AND all Kogenate royalties

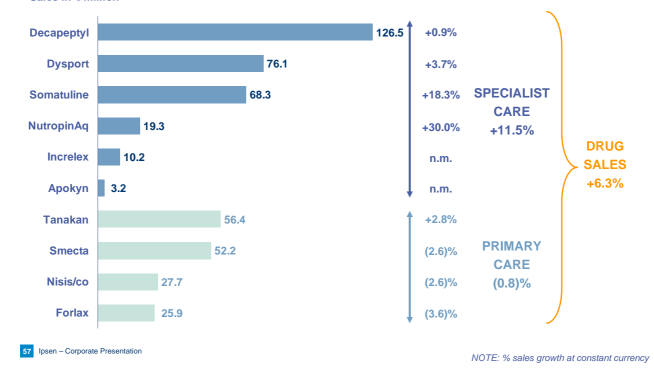
First half 2009 detailed financial performance

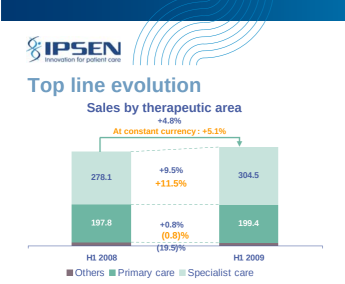


Main products performances

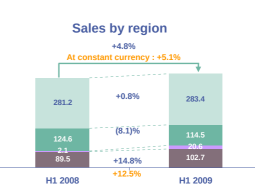
Sales in € million

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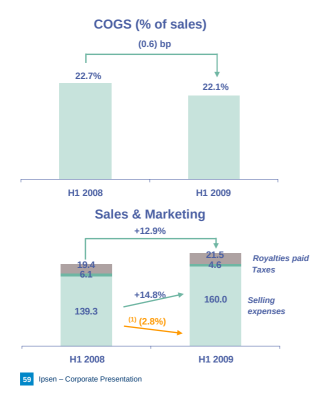






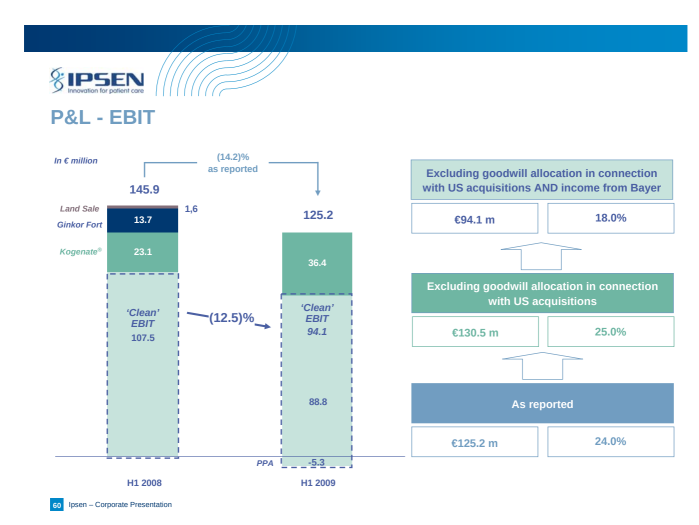






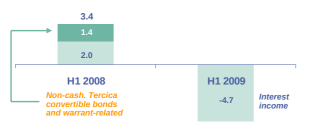


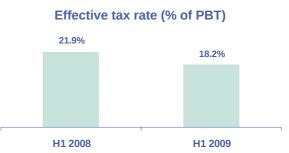
NOTE 1: excluding US



P&L – below EBIT

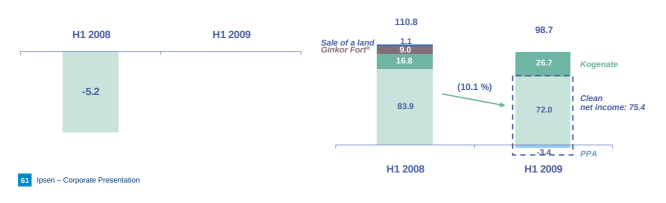






Consolidated result (€m - group share)

Income from Associates (€m)



SIPSEN Innovation for patient care

Balance	Sheet	evo	lution
Balanoo			

- In million of euros Asset	s		- In million of euros	ties	
	31 Dec 08	30 Jun 09		31 Dec 08	30 Jun 09
Goodwill	^(*) 290.8	290.8	Equity	(*) 885.0	928.4
Property. plans & equipments	237.9	244.7	Minority interests	1.6	1.8
Intangible assets	(*) 232.9	239.3	Total equity	(*) 886.6	930.2
Other non-current assets	(*) 112,9	140.5	Long-term financial debts	160,4	12.7
Total non-current assets	(*) 874.5	915.3	Other non-current liabilities	(*) 196,4	276.3
Total current assets	(*) 688.6	589.7	Short-term debts	8.3	8.0
Incl. cash and cash equivalents	239.6	140.2	Other current liabilities	307.8	275.1
Assets / discontinued operations	1.3	0.7	Liabilities / discontinued operations	4.9	3.5
Total assets	1 564,4	1 505.8	Total Liabilities	1 564,4	1 505.8
Net Cash	66.2	118.9			



Cash flow statement

	30 Jun 08	30 Jun 09	Comments
- In million of euros			L. Deferred more not in second
Cash Flow before change in working capital	141.3	121.5	 Deferred revenues net increase: €+56.7m (Medicis / Galderma)
- Increase / Decrease in working capital	(17.1)	25.7	
Net cash flow generated by operating activities	124.1	147.2	 Receivables, payables, inventory and others: €-31.0m
Investment in intangible assets and property. plant & equipment	(34.2)	(25.6)	
Others	1.8	(6.8)	Tangible assets: €-14.7m
Net cash flow used in investing activities	(32.4)	(32.4)	 Intangible assets: €-10.9m
Net change in borrowings	(9.8)	(159.4)	
Dividends paid	(55.2)	(58.2)	\backslash
Others	0.1	-	reimbursement of credit facility:
Net cash flow used in financing activities	(64.9)	(217.6)	€-150m
Discontinued operations	(1.0)	(0.2)	 Shares buy back: €-6m
Change in cash and cash equivalent	25.8	(103.0)	
Impact of exchange rate fluctuations	(3.0)	4.8	
Closing cash & cash equivalents	263.7	139.1	
Closing Net Cash	239.4	118.9	

Financial appendices





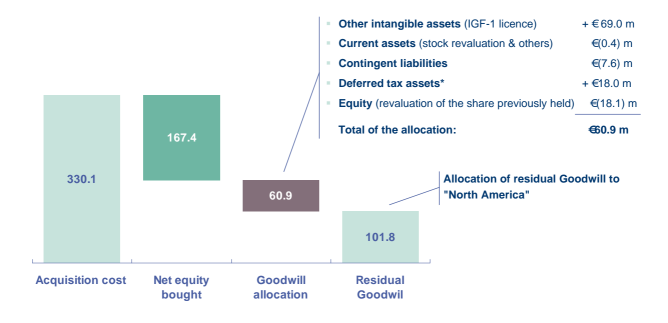
Cash flow generation



borrowings and other financial liabilities plus or minus derivative financial instruments



Allocation of the Tercica purchase price accounting





Milestones Cashed in but not yet Recognised as Revenues

- In million of euros	30 Jun 08	30 Jun 09
Payments recognised as revenues in year N+1	11.2	12.1
Payments recognised as revenues in years N+2 and beyond	205.7	195.2
Total Milestones cashed in but not yet recognised as revenues	216.9	207.3



Decrease linked to global consolidation of Tercica and elimination of deferred revenues on Somatuline US