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MOXDUO®

Morphine + Oxycodone



QRxPharma Snapshot

- Australian based pain drug developer with offices in Sydney and Bedminster, NJ (ASX:QRX, OTCQX:QRXPY)
- Comprehensive portfolio around MOXDUO product line
- Blockbuster potential: global opioid market estimated at \$14 billion annually¹
- Strategic commercialisation collaborations with Actavis Inc. (US) and Paladin Labs Inc. (Canada)
- MOXDUO Complete Response Letter (CRL) issued by the U.S. FDA in June 2012; Productive Review Meeting in August 2012
 - Additional data package to FDA 2H, 2012; NDA re-filing with FDA decision anticipated around mid-2013



Solid Foundation for Growth

- Multiple formulations for complete "hospital-to-home" care; expected patent exclusivity through 2029
- MOXDUO delivers equal or better pain relief with fewer side effects than current treatments
- Potential changes in US regulatory policy are favourable for commercialisation of MOXDUO IR
- Double digit royalties on US sales of MOXDUO IR
- Experienced management team and board of directors
- KOL confidence in MOXDUO IR as a potential therapeutic option; Company commitment to bringing product to market



MOXDUO Product Portfolio From Hospital to Home

	MOXDUO® IR	MOXDUO® CR	MOXDUO [®] IV
Delivery	Immediate Release	Controlled Release	Intravenous
Target	Moderate to severe acute pain	Chronic pain (i.e. osteoarthritis, back, neuropathic)	Hospital based: moderate to severe acute pain
Formulation	Oral Capsule	Oral tablet w/abuse deterrent	Injectable
Partnerships	Actavis, Inc. and Paladin Labs Inc. US and Canada commercialisation		
Status	US: Additional Data Package and NDA re-filing Canada: NDS application expected to lodge 1H, 2013	Phase 1 Complete	Phase 2 Formulation development



MOXDUO®

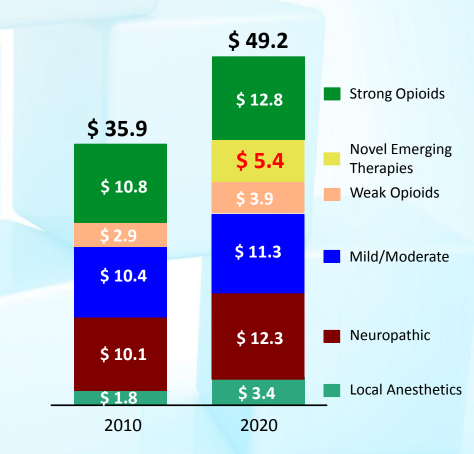
THE PAIN MARKET



Current Global Pain Market

Drug Class Sales for Pain in Major Pharmaceutical Markets, 2010 – 2020 (US\$ billions) ¹

- Large market opportunity: US\$14 billion¹
- Opioids are the "gold standard"
- Limited product innovation
- Strong opioids forecasted to maintain sales dominance through 2020 (aging population)
- Payors and KOLs: "need for better pain relief with fewer side effects"





Acute Pain Market AT-A-GLANCE



US Market Opportunity

- \$2.5B US Market with 210M+ annual Rxs (CAGR of 5-6%)
- Acute pain affects 75M Americans
- Limited product innovation; regulatory hurdles for new therapies
- Limited branded competition expected near-term



Clinical Unmet Needs

- Inadequate postoperative pain management
- Reduction in opioid-related AEs, specifically GI & CNS that limit their use
- Existing acute pain drugs associated with hepatic and GI toxicities

In the US, 6+ prescriptions are written for an acute opioid every second.



Proposed U.S. Regulatory Changes

2011 2012 2013 2014



FDA vote to reduce acetaminophen in combination products (i.e. Vicodin/Percocet)

FDA issues mandate to reduce acetaminophen to < =325 mg

Bipartisan Bill in CMTE to reschedule hydrocodone



Potential enactment of hydrocodone legislation; harder to prescribe Vicodin



Mandated time for withdrawal of combo products containing >325mg

- FDA mandated lower strength opioid/APAP combos will decrease efficacy and increase number of patients needing acute pain medicine
- Potential rescheduling of Vicodin to Schedule 2 will make it harder to prescribe and decrease number of prescriptions
- Creates void of about 50% of acute pain market (100 million Rx's)



Disrupted Acute Pain Market Provides MOXDUO IR Opportunity

2010 US Prescription Market Share for Acute Pain Opioids





Source: IMS 2010



IMMEDIATE RELEASE



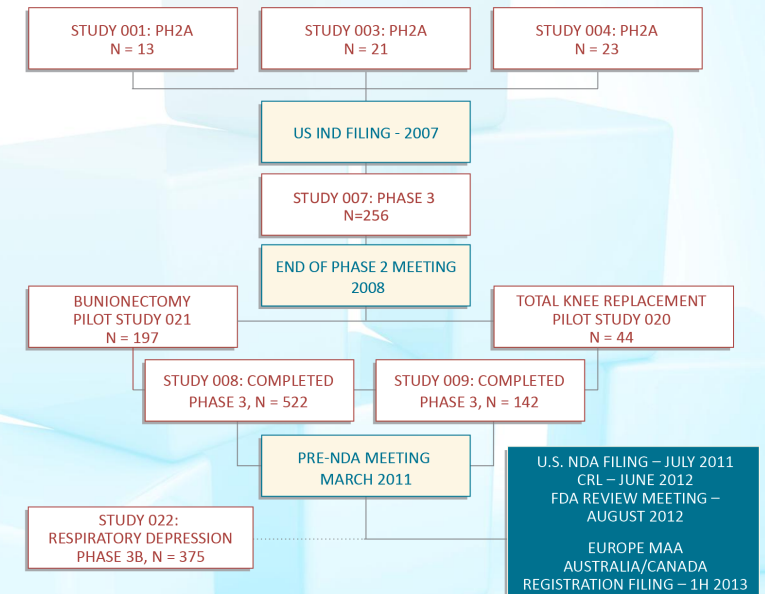
MOXDUO IR Product Profile

Key Features

- First line therapy for the treatment of moderate to severe acute pain
- Only opioid-opioid combination product available
- Immediate release formulation of morphine and oxycodone in a fixed3:2 ratio in capsules of the strengths:
 - 3 mg/2 mg
 - 6 mg/4 mg
 - 9 mg/6 mg
 - 12 mg/8 mg
- Four to six hourly dosing
- Demonstrated reduction in the occurrence and intensity of clinically significant opioid-related side effects compared to morphine, oxycodone and Percocet®



MOXDUO IR Clinical Development Path





Key Trial Conclusions

- Bunionectomy Trials: Pilot 021 & Pivotal 008 (n=719)
 - Met primary analgesic efficacy endpoint vs. morphine and oxycodone
 - MOXDUO IR proven superior to components on efficacy measures
 - Consistent safety advantage of MOXDUO IR
 - Pilot: 50-75% lower frequency of moderate to severe nausea, vomiting & dizziness compared to equi-analgesic doses of morphine or oxycodone
 - Phase 3: Despite higher dose and better pain relief of MOXDUO than morphine or oxycodone, AE rate and duration not statistically different

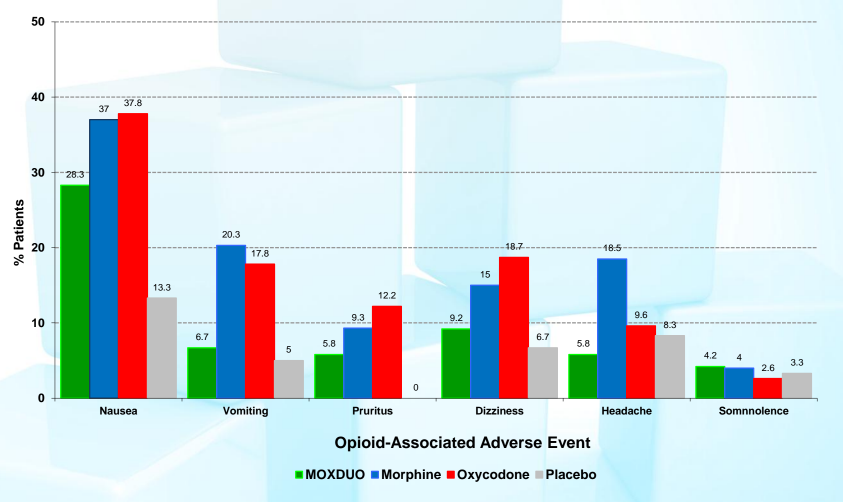


Key Trial Conclusions

- Total Knee Replacement Trials: Pilot 020 & Pivotal 009 (n=186)
 - Met all primary analgesic efficacy endpoints vs. Percocet
 - Pilot: MOXDUO superior to Percocet
 - Pivotal: MOXDUO high dose better pain relief than low dose
 - Frequency of AEs much lower than Percocet
 - Significant pharmacoeconomic benefit: improved time to walk, sleep, etc.



Equi-analgesic doses of MOXDUO, Morphine, Oxycodone, vs. Placebo – All Studies







Exploratory Phase 3 Study 022

Respiratory Depression Study

Objectives

- Europe: Support MAA as per 2010 Scientific Advice Meeting with the BfArM; comparative AE labeling; respiratory safety advantage; overall risk / benefit
- US: Help prepare for future definitive studies for comparative AE information in PI
- **US NDA**: Provide important opioid safety information regarding respiratory function advantages of MoxDuo relative to equi-analgesic doses of morphine and of oxycodone

Study Design and Outcome			
Phase	Phase 3		
N	375		
US Sites	6		
Design	Randomized 1: 1: 1, double-blind, multicenter, repeat dose, 3 arms stratified by gender and by age (≥ 60 yrs or < 60 yrs; 40% of patients were age 60+)		
Doses / Schedule	MoxDuo IR 12 mg/8 mg vs. Morphine 24 mg vs. Oxycodone 16 mg Every 6 hours for 48 hours		
Primary Endpoints	To explore the effects of MoxDuo IR relative to morphine and oxycodone comparators on oxygen desaturation, a measure of respiratory impairment		
Secondary Endpoints	Percent of subjects with moderate or severe, spontaneously reported, treatment emergent events of nausea, emesis, or dizziness		

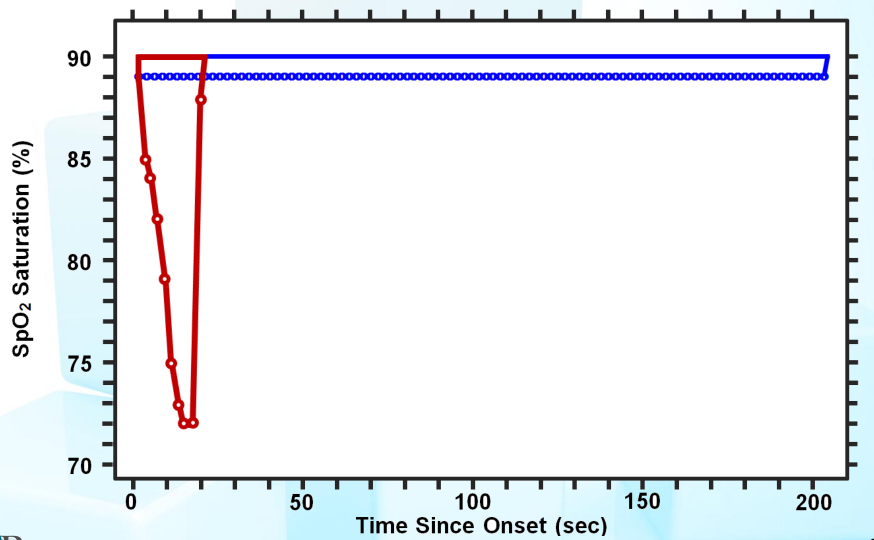


Effects of Opioids on Blood Oxygen Levels

- Respiratory depression is enhanced by opioids
- Death from opioid overdose is due to respiratory depression
- Pulse oximetry continuously monitors blood oxygen levels
 (SpO₂) using finger tip sensor
- SpO₂ normal values 96-100%; <90% = oxygen desaturation
- This study used electronic records of SpO₂ values
- SpO₂ desaturations (intensity and incidence) are key endpoints

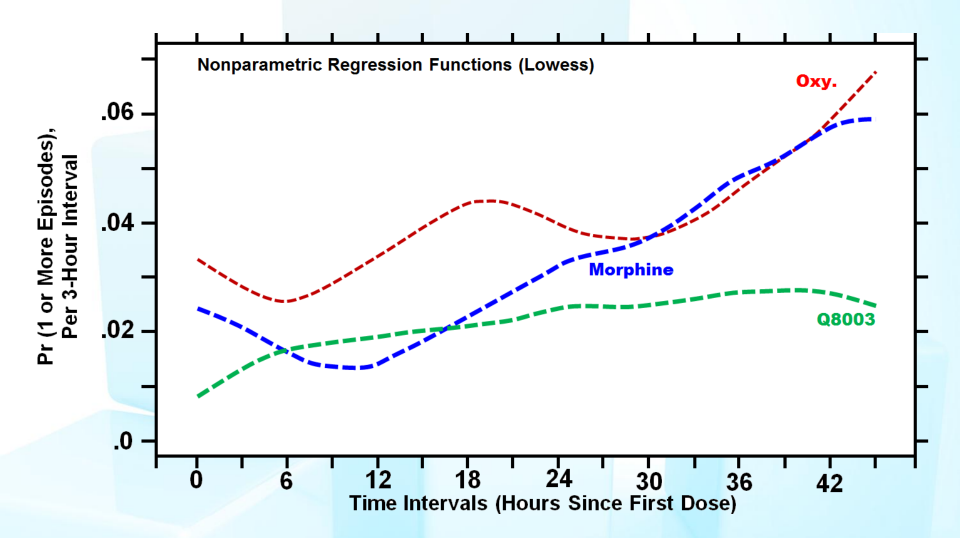


Example of actual curves of two different patterns of SpO₂ desaturations



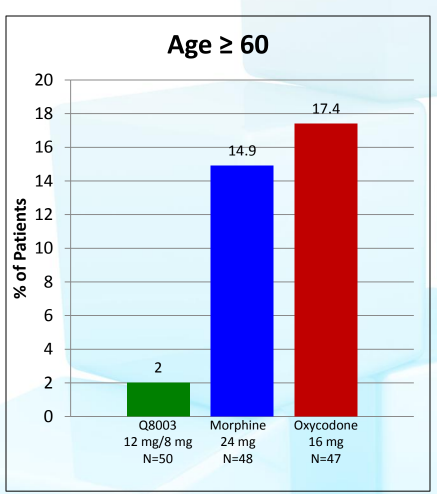


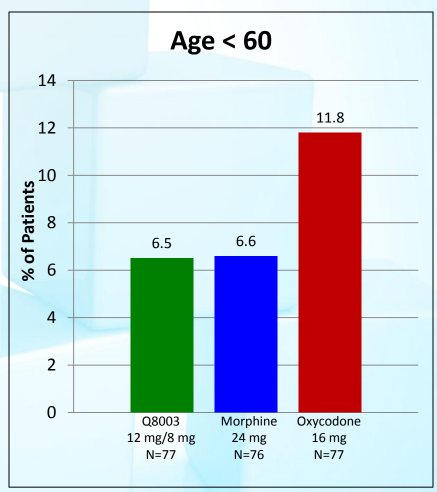
Lowess Curves at 80% SpO₂ Severity Cut Point





Percent of Subjects with Sustained (30+ sec.) SpO₂ Desaturations by Age at 70% Cut Point







Respiratory Safety Conclusions

- Oxygen desaturation data demonstrate a beneficial safety signal relative to equi-analgesic doses of morphine and of oxycodone
- The risk of a patient experiencing the more medically meaningful desaturations is appreciably less for MOXDUO IR than for morphine or oxycodone treatments
- Lower likelihood of experiencing severe desaturations over time may reduce the possibility of dangerous respiratory morbidity when patients are discharged from the hospital
- Desaturation data from older patients (≥ 60 y/o) provides evidence of a respiratory safety benefit for MOXDUO IR in an important patient subpopulation



Conclusions

- Over the entire NDA program, patients receiving morphine and oxycodone were 9%-96% more likely to experience an opioid-like adverse event than MOXDUO IR treated patients
- These differences in favor of MOXDUO IR were seen for the following TEAES: nausea, vomiting, dizziness, pruritus, headache, somnolence/sedation and decreased oxygen desaturation when directly compared to equal-analgesic doses of either morphine or oxycodone



MOXDUO Safety Advantage

STUDIES OF COMBINED OPIOIDS CONSISTENTLY PROVIDE EQUIVALENT EFFICACY WITH A SIGNIFICANT REDUCTION IN OPIOID-RELATED MODERATE TO SEVERE ADVERSE EVENTS

QRx STUDIES

MOXDUO IR Study 022 (bunionectomy)	Oxygen desaturation less severe and of shorter duration compared to equianalgesic doses of Morphine or Oxycodone
MOXDUO IR Study 021 (bunionectomy)	50-75% reduction in moderate to severe nausea, vomiting and dizziness compared to equianalgesic doses of Morphine and Oxycodone
MOXDUO IR Study 020 (knee replacement)	100% reduction in moderate to severe nausea and emesis compared to the Percocet
MOXDUO IV Study (hip replacement)	35% reduction in nausea and 38% reduction in emesis Compared to IV Morphine
MOXDUO Two Phase 2 trials in Australia (chronic pain)	34-40% decrease in the amount of drug to achieve equianalgesia compared to oral morphine. Decreased rate of drowsiness, dizziness, constipation and nausea.

INDEPENDENT STUDIES¹

Blumenthal et al 2007 (Spinal discectomy)	>	80-100% reduction in nausea and emesis compared to PCA Morphine
Jamison et al 1998 (Chronic low back pain)	>	17-49% reduction in intensity of a range of adverse events compared to Oxycodone
Lauretti et al 2004 (Cancer pain)	>	86% reduction in nausea and 100% reduction in emesis compared to Morphine.



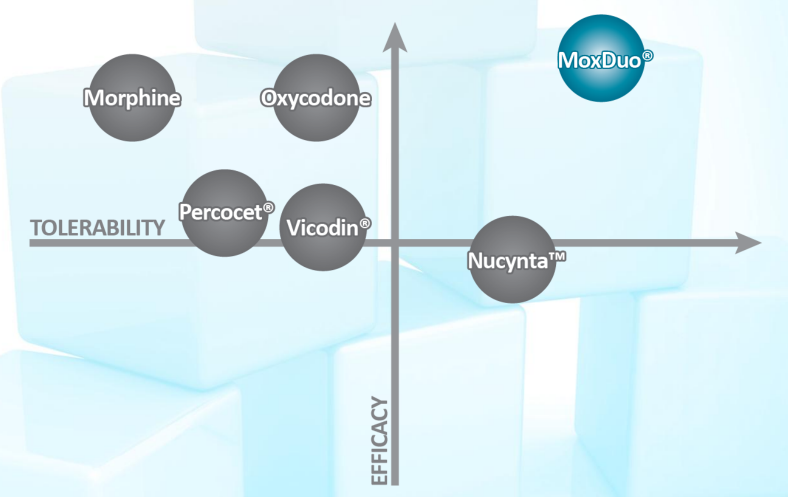
MOXDUO Pharmacoeconomic Benefits

- Versus Percocet® TKR study, MOXDUO IR patients were out of bed faster, walked and slept better
- US study finds \$4,880 \$36,152¹ incremental costs in patients suffering GI side effects following treatment with IR opioids
 - Extended hospitalization, increased nursing care and re-admissions
- Reimbursers, managed care and KOL feedback
 - Significant benefit from decreasing hospitalization by as little as 4 hours or recovery room time by 20 minutes

MOXDUO's advantageous safety profile may improve patient recovery, decrease hospital time and lower total cost of care.



MOXDUO IR Value Proposition = Greater Tolerability + Equal/Better Analgesia





MOXDUO®

Immediate Release Commercialisation Plan



Actavis Strategic Partnership

- Exclusive US commercialisation and development rights for MOXDUO IR
 - Actavis pays all product launch, marketing and sales costs
- 10%-30% royalties based on net sales thresholds from launch
 - Except 50% royalty on \$150m of cumulative sales (starting from 3-6 months following product launch)
- QRxPharma retains a right to co-promotion/profit-share
 - Option to create sales force and provide up to 25% of the effective selling effort to US prescribers after first 12 months of launch
- QRxPharma retains ownership of MOXDUO IR outside the US and Canada





- European based Actavis Group hf is a leading manufacturer of branded and generic opioids worldwide
 - Revenues of ~EUR 1.8 billion & 10,000 employees worldwide
 - US sales highlighted by Kadian® (extended release morphine)
- Watson Pharmaceuticals acquisition proposed in April 2012; completed in November 2012
 - Creates 3rd largest global generics company
 - ~\$8.0 billion projected 2012 pro forma combined revenue
 - Maintaining commitment to branded marketplace
- MOXDUO IR commercialisation preparation ongoing





- Strategic collaboration with Paladin Labs for Canadian commercialisation rights of immediate release MOXDUO
- QRxPharma to receive double-digit royalties and up to US\$25M in milestone payments on sales; upfront payment of US\$500,000
- Paladin Labs pays all regulatory, product launch, marketing and sales costs
- QRxPharma retains Canadian rights to MOXDUO IV and CR
- Paladin Labs is a leading specialty pharmaceutical company based in Montreal and listed on the Toronto Stock Exchange
 - Branded pain products include: Metadol[®]; Pennsaid[®]; Tridural[®]; and Abstral[®]



Mitigating Reimbursement Risk

- Appropriate pricing, contracting and patient pull-through
- Advantaged market other acute pain products (Vicodin® and Percocet®)
 being reduced due to safety issues and potential rescheduling
- Potential for significant pharmacoeconomic benefits recognized by payers/KOLs
- Reimbursement strategy = Tier 3 Formulary
 - Insurance companies will not have to pay more
 - Customer co-pays are manageable



MOXDUO®

Pipeline Potential



MOXDUO CR (Controlled Release)

- Sustained release formulation to provide at least 12 hours of analgesia for moderate to severe chronic pain
- Abuse deterrent and tamper resistant features
- Phase 1 results showed:
 - High bioavailability and complete absorption
 - One fifth the variability of OxyContin; will provide very stable plasma levels when given twice daily
 - Lower peaks and higher troughs should lead to better safety & lower side effects; better tolerability at higher doses
 - Should be an effective once or twice daily treatment
- Current formulation will progress to Phase 2



MOXDUO: Peak Sales Potential

		MOXDUO IR	MOXDUO CR	MOXDUO IV
Market Size		■ ~200 mm Rx (2012)	■ ~34 mm Rx (2015)	■ ~29 mm Rx (2014)¹
		■ Annual market growth 1.0%	■ Annual market growth 3.0%	■ Annual market growth 1.0%
		■ QRx targets ~ 50% of market	■ QRx targets 100% of market	QRx targets 100% of market
Market		■ Initial share: 1.0% (2012)	■ Initial share: 1.4% (2015)	■ Initial share: 1.5% (2014)
Penetrati	on	■ Peak share: 5.0% (2015)	■ Peak share: 13.9% (2020)	■ Peak share: 13.0% (2018)
Pricing		 Initial price: \$112 based on 4 doses per day and 14 days of therapy 	Initial Rx Price: \$180 based on 2 doses per day and 30 days of therapy	Initial price: \$32 based on 4 vials per day and 2 days of therapy
		■ Annual price increase: 5.0%	■ Annual price increase: 5.0%	■ Annual price increase: 5.0%
		■ Peak sales: ~\$680 mm	■ Peak net sales: ~\$1,300 mm	■ Peak net sales: ~\$150 mm
Blockbus Opportur		■ Paracetamol Limitation -Peak sales: ~\$1,350 mm	Oxycontin - \$3 billion/year - off patent in 2013, opening market	
		■ plus Vicodin Rescheduling - Peak sales: ~\$2,000 mm	for MOXDUO CR in 2015	



1 Rx represents "eaches".

MOXDUO®

Company Overview



Leadership Team

Senior Management

- John Holaday, PhD (CEO)
- Ed Rudnic, PhD (COO)
- Chris Campbell (CFO)
- Richard Paul, MD (EVP Drug Development)
- Warren Stern, PhD (Clinical Consultant)
- Janette Dixon, PhD (VP Global BD)
- Patricia Richards, MD, PhD (CMO)

Board of Directors

- Peter Farrell, PhD Chairman (ResMed)
- Michael Quinn (Innovation Capital)
- Peter Campbell (Sonic Healthcare)
- Gary Pace, PhD (ResMed, founder QRxPharma)
- John Holaday, PhD (CEO)

Scientific Advisory Board

- Solomon Snyder, MD (Chair)
- Lester Crawford, DVM, PhD
- Robert Lenox, MD
- Michael J Cousins, MD, AM
- Horace H Loh, PhD
- Gavril Pasternak, MD, PhD
- · Richard Payne, MD



Financial Summary (10 December 2012)

Shares on issue: 145 million (ordinary)

Market cap: A\$106 million

Cash on hand:

30 September 2012 A\$18.7 million (last reported)

Cash burn: CY2013

Share registry: +80% institutional / HNW

Listing: ASX: QRX / OTCQX: QRXPY



MOXDUO IR Key Milestones

DATE	MILESTONE
✓ July 2011	NDA submission to FDA
✓ December 2011	Signed strategic collaboration with Actavis
× 25 June 2012	NDA PDUFA Date; CRL Received
✓ August 2012	FDA Review Meeting
✓ October 2012	Signed strategic collaboration with Paladin
• 2H, 2012	Submission of additional data package requested by FDA
• Mid-2013	Anticipated decision from FDA on a refiled MOXDUO NDA
• 1H, 2013	Submit additional regulatory filings: Europe, Australia & Canada
• 2H, 2013	Product launch in the US



Investment Highlights

- Comprehensive Portfolio: MOXDUO delivers equal or better pain relief with fewer side effects than current treatments
- Actavis commercialisation partnership in the US; Paladin Labs
 commercialisation partnership in Canada: MOXDUO IR to be key branded
 pain product for Actavis / Watson
- Advantaged market: Favourable US regulatory and potential prescription scheduling changes
- Blockbuster potential: Global opioid market estimated at \$US14bn¹
- Strong IP: Expected patent exclusivity through 2029
- Expanded pipeline: Further progress MOXDUO CR and MOXDUO IV products



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