

Resverlogix BET Inhibition for Global Vascular Risk BIO CEO & Investor Conference New York, NY

February 12-13, 2018

Forward Looking Statements

RESVERLOGIX

This presentation may contain certain forward-looking information as defined under applicable Canadian securities legislation, that are not based on historical fact, including without limitation statements containing the words "believes", "anticipates", "plans", "intends", "will", "should", "expects", "continue", "estimate", "forecasts" and other similar expressions. In particular, this presentation includes forward looking information relating to the Company's clinical trials and the potential role of apabetalone in the treatment of CVD, DM, chronic kidney disease, Orphan diseases, and peripheral artery disease. Our actual results, events or developments could be materially different from those expressed or implied by these forwardlooking statements. We can give no assurance that any of the events or expectations will occur or be realized. By their nature, forward-looking statements are subject to numerous assumptions and risk factors including those discussed in our Annual Information Form and most recent MD&A which are incorporated herein by reference and are available through SEDAR at www.sedar.com. The forwardlooking statements contained in this presentation are expressly qualified by this cautionary statement and are made as of the date hereof. The Company disclaims any intention and has no obligation or responsibility, except as required by law, to update or revise any forward-looking statements, whether as a result of new information, future events or otherwise.

Main Subject Matter	 Apabetalone, the only-in-class BET inhibitor (BETi), reduces key risk factors for high risk cardiovascular and renal patients resulting in reduction of Major Adverse Cardiac Events (MACE), observed renal improvement markers
Advanced Mechanism	 Epigenetic modulation of gene expression makes BETi a novel approach No known BETi competitor for next 9 plus years
Confirmed Science	 Proteomics, genomics, pathway analysis, mechanism of action are all very well understood
Clinical Evidence	 Phase 2b data – up to 62% RRR of MACE in high risk CVD patients Phase 3 BETonMACE trial 90% enrolled CVD/CKD risk biomarkers tracked to date - positive
Corporate Expansion	Resverlogix corporate goal is to expand commercial partner program

Capitalization and Financial Profile

Founded	2001		
Ticker	TSX: RVX		
Market Cap	~C\$300MM		
Long Term Debt	~C\$0.0MM		
Shares Outstand	175.04MM		
Cash Burn (Annual)	~C\$40.0M		
Finance	\$87MM – Announced October 2017		

RVX Top Shareholders



- RVX shareholder base consists of several long term investors who have been supportive over 10 years
- RVX maintains a diversified public market float of approximately 54M shares or ~\$130MM

2017 Major Accomplishments



Upcoming Clinical Year Estimates



Apabetalone in the Clinic



Apabetalone has been tested in multiple clinical trials with a good safety and efficacy profile

BET Literature Impact Growing: CVD and Renal Risk



OPEN OACCESS Freely available online				
RVX-208, an Inducer of ApoA-I in I Bromodomain Antagonist		DOI 10 ORI	Cardiovase Drugs 0.1007/s40256-017-0250-3 GINAL RESEARCH ARTICLE Ective BET Protein Inhibition with Apab	CrossMark
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Sylwia Wasiak ^a , Dean Gill Christopher Halliday ^a , Kai Kevin G. McLure ^a , Peter R Ewelina Kulikowski ^a , Jan Norman C. Wong ^{a,*} ^a Resverlogix Corp., Calgary, Canada ^b Resverlogix Corp., San Francisco, USA	< Previous Article Article in Press	Α	Inticles in Press Next Article >	otic cardiovascular ys that contribute to CrossMark
J. of Cardiovase. Trans. Res. DOI 10.1007/s12265-017-9755-z ORIGINAL ARTICLE	Benefit of Apabetalor		Plasma Proteins in Renal Disease	sson ^b , Michael Sweeney ^b ,
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Sylwia Wasiak ¹ • Dean Gilham ¹ • Laura M. Tsujikawa ¹ • Christopher Halliday ¹ • Cyrus Calosing ¹ • Ravi Jahagirdar ¹ • Jan Johansson ² • Michael Sweeney ² • Norman C. Wong ¹ • Ewelina Kulikowski ¹			Ravi Jahagirdar, Sarah Attwell, Suzana Marusic, Alisor Kevin G. McLure, Dean Gilham, Karen Norek, Henrik Jennifer Tobin, Gregory S. Wagner, Peter R. Young, I and Ewelina Kulikowski	C. Hansen, Raymond Yu,
Received: 21 December 2016 / Accepted: 17 May 2017 © The Author(s) 2017. This article is an open access publication TSX: RVX	1	RES	Resverlogix Corporation, Calgary, Alberta, Canada (R.J., S.A., K.G.M., D.G., K E.K.); Hooke Laboratories Inc., Lawrence, Massachusetts (S.M.); Bolder BioP Inc., Sunnyvale, California (N.S.)	

Unique Mechanism of Action



BET Inhibition Impacts the Pathways that Drive Cardiovascular Disease and Kidney Diseases



Apabetalone, a bromodomain extra-terminal (BET) protein inhibitor, inhibits BRD4, thereby regulating the expression of genes and restoring the function of pathways underlying the pathogenesis of CVD and kidney disease



Nicholls et al. 2017: American Journal of Cardiovascular Drugs





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MACE: <u>Major Adverse Cardiac Events including: death,</u> myocardial infarction, stroke, coronary revascularization, hospitalization for acute coronary syndrome or heart failure

Decrease in MACE was most profound in patients who had a higher level of inflammation such as patients with diabetes

BETonMACE CV Outcomes Study Design





The study is an event-based trial and continues until 250 narrowly defined MACE events have occurred



Primary Objective

To evaluate if treatment with apabetalone as compared to placebo increases time to the first occurrence of triple MACE. Triple MACE is defined as a single composite endpoint of: 1) CV death or 2) non-fatal MI or 3) stroke.

Key inclusion criteria

- Type II Diabetes Mellitus
 HbA1c > 6.5% or history of diabetes medications
- CAD event 7 days 90 days prior to screening
 Myocardial infarction (MI), unstable angina or percutaneous coronary intervention
- HDL <40 mg/dL for males and <45 mg/dL for females

Primary Endpoint

Time from randomization to the first occurrence of adjudication-confirmed triple MACE defined as a single composite endpoint of: 1) CV Death or 2) Non-fatal MI or 3) Stroke.

Secondary Endpoint

- Time from randomization to the first occurrence of adjudication-confirmed MACE including revascularization and unstable angina
- Changes in apoA-I, apoB, LDL-C, HDL-C, and TG
- Changes in HbA1c, fasting glucose, and fasting insulin
- Changes in ALP and eGFR

Screening & Baseline Clinical Chemistry As of December 4, 2017



Parameter	Ν	Median (min, max)	
Age	2,091	62 (33, 88)	
Alkaline Phosphatase [†] , U/L	2,065	78 (5, 915)	
HDL-C, mg/dL	2,074	33 (14, 47)	
hsCRP [†] , mg/L	425	2.9 (0.2, 162.1)	
Fibrinogen [‡] , mg/L	406	387 (92, 730)	
LDL-C, mg/dL	2,057	65 (3, 232)	
Apolipoprotein A-I [†] , mg/dL	415	118 (58, 179)	
Glucose, mg/dL	2,074	135 (41, 555)	
HbA1c, %	2,035	7.3 (4.5, 15.1)	
Platelets, 10 ⁹ / L	1,976	248 (6, 989)	
NLR, ratio	1,993	2.6 (0.6, 16.5)	
Males	75.6% males		
Statin Allocation	52% atorvastatin	48% rosuvastatin	
+ results from visit 2/wk 0, whereas all other values are from visit 1/scree			

† results from visit 2/wk 0, whereas all other values are from visit 1/screening



- ~ 90% enrolled
- CKD Subgroup: ~11% of patients have eGFR<60 at screening
- Cognition Subgroup: ~18% of patients have completed MoCA at Baseline; Target patients are those with baseline MoCA ≤ 25
- Consistent data repeatable positive effects in key CVD and CKD biomarkers
- New data to target MoCA in elderly cognition subgroup (70 and over)
- Projected primary MACE rate still 8.0 per 100 patient years on top of aggressive standard of care = strong unmet need

Patient Enrichment Strategy



Sources: Calculated from CDC Heat Disease Facts; Holden, SE. et al. 2015; White, WB. et al. 2013; Kim, H. et al. 2015; Cardarelli, F. et al. 2008; Okada, T. et al. 2008

Market Opportunity Pathways





• 6 Million Patients

Sub-population data analysis
 Cognition and dementia
 2+ Million Patients

Improving Global Vascular Risk

Balanced for Success!



THREE KEY DEVELOPMENT TARGETS ARE IN PLACE

INNOVATION

Resverlogix owns the worlds most advanced BRD4 epigenetics program

EFFICACY Clinical and safety data is continuing to suggest a successful program

PHARMACO-ECONOMICS

The payor groups now hold the power to determine success

Based on SUSTAIN/ASSURE, NNT and cost/event prevented were favourable versus comparators in their respective patient populations



Note: Medication cost based on US WAC cost (PriceRx); Kaplan Meier estimates are based on digitized curves from publications ^No patient population standardization has been applied so additional population specific factors beyond severity can also influence NNT values *1 year showed no benefit to calculate NNT; estimated by taking 5 year KM rate of 50 x 5 years

Apabetalone HEOR Evidence Benchmarking - Final Results v4.0 Oct 2015

imshealth brogan

RVX testing further price bands: Tier 3 based on higher risk populations

Pricing & Reimbursement: V2 Report 2016



- RVX performed multiple outreach reports with leading US KOL payers for market pricing analytics
- Apabetalone target plan: higher risk CVD patients (e.g. Diabetes with recent ACS, CKD, Dialysis, Dementia) supported positive pricing and reimbursement with leading US payer groups
- Higher risk patients represent significantly increased burdens to healthcare systems on account of greater costs associated per patient per year
- Payer responses shows strong support for pricing value proposition falls within ICER range of \$140-175K USD. This ICER range represents superior value proposition versus current CVD risk competitors such as PCSK9s and SGLT2s
- Global pricing band planned by US market first, then European, Canada with applicable discounts

Payer KOL Outreach: Key Payer Support



Organization	Lives Covered	MACE Reduction: Unmet need in Recent ACS and T2DM patients	MACE Reduction: Unmet need in CKD patients	ICER Threshold per annum
Payer 1	55 M	Moderate to High	Moderate to High	\$ < 100,000
Payer 2	65 M	Moderate to High	Moderate to High	\$ < 200,000
Payer 3	37 M	Moderate to High	Moderate to High	\$ < 100,000
Payer 4	40 M	Moderate to High	Moderate to High	\$ < 150,000
Payer 5	11 M	Moderate to High	Moderate to High	\$ < 150,000

- 5 Payers 208 million lives covered, Key C Suite executives contacts President, Chief Medical Directors, COO, Executive VP Pharmacy
- Pricing bands support average ICER Target Threshold of approximately **\$140,000-\$200,000** USD
- Pricing bands support average price of **\$6,000 \$12,000** based on new enriched high risk patients

Apabetalone Opportunity

Highlights

- Novel, first in class, technology no competitor 8 10 years
- Clear science and clinical data supporting strong rationale for risk reduction
- Growing BET literature publications in CVD / Renal risk
- Strong KOL Payers and Prescriber support
- Transformative science and unprecedented commercial opportunity