
**UNITED STATES
SECURITIES AND EXCHANGE COMMISSION**
Washington, D.C. 20549

FORM 8-K

**CURRENT REPORT
Pursuant to Section 13 or 15(d)
of the Securities Exchange Act of 1934**

Date of Report (Date of earliest event reported): **January 4, 2019**

RHYTHM PHARMACEUTICALS, INC.
(Exact name of registrant as specified in its charter)

Delaware
(State or other jurisdiction
of incorporation)

001-38223
(Commission
File Number)

46-2159271
(IRS Employer
Identification Number)

500 Boylston Street, 11th Floor
Boston, MA 02116
(Address of principal executive offices, including Zip Code)

Registrant's telephone number, including area code: **(857) 264-4280**

Check the appropriate box below if the Form 8-K filing is intended to simultaneously satisfy the filing obligation of the registrant under any of the following provisions:

- Written communications pursuant to Rule 425 under the Securities Act (17 CFR 230.425)
- Soliciting material pursuant to Rule 14a-12 under the Exchange Act (17 CFR 240.14a-12)
- Pre-commencement communications pursuant to Rule 14d-2(b) under the Exchange Act (17 CFR 240.14d-2(b))
- Pre-commencement communications pursuant to Rule 13e-4(c) under the Exchange Act (17 CFR 240.13e-4(c))

Indicate by check mark whether the registrant is an emerging growth company as defined in Rule 405 of the Securities Act of 1933 (§230.405 of this chapter) or Rule 12b-2 of the Securities Exchange Act of 1934 (§240.12b-2 of this chapter). Emerging growth company

If an emerging growth company, indicate by check mark if the registrant has elected not to use the extended transition period for complying with any new or revised financial accounting standards provided pursuant to Section 13(a) of the Exchange Act.

Item 7.01. Regulation FD Disclosure.

On January 4, 2019, Rhythm Pharmaceuticals, Inc. (the “Company”) issued the attached press release regarding updated clinical data and expected 2019 milestones to be presented at the 37th Annual J.P. Morgan Healthcare Conference on Thursday, January 10, 2019. Copies of the press release and the presentation slides are furnished as Exhibits 99.1 and 99.2, respectively, to this Current Report on Form 8-K. The Company undertakes no obligation to update, supplement or amend the materials attached hereto.

The information contained in this Item 7.01 and in the accompanying Exhibits 99.1 and 99.2 shall not be incorporated by reference into any filing of the Company, whether made before or after the date hereof, regardless of any general incorporation language in such filing, unless expressly incorporated by specific reference to such filing. The information in this Item 7.01 and the exhibits hereto shall not be deemed to be “filed” for purposes of Section 18 of the Securities Exchange Act of 1934, as amended, or otherwise subject to the liabilities of that section or Sections 11 and 12(a)(2) of the Securities Act of 1933, as amended.

Item 9.01 Financial Statements and Exhibits.

<u>Exhibit No.</u>	<u>Description</u>
99.1	Press release dated January 4, 2019.
99.2	Company Presentation dated January 2019

SIGNATURES

Pursuant to the requirements of the Securities Exchange Act of 1934, as amended, the registrant has duly caused this report to be signed on its behalf by the undersigned hereunto duly authorized.

RHYTHM PHARMACEUTICALS, INC.

Date: January 7, 2019

By: /s/ Hunter Smith
Hunter Smith
Chief Financial Officer



Rhythm Pharmaceuticals to Outline Expected 2019 Milestones and Review Updated Clinical Data in Bardet-Biedl Syndrome at 37th Annual J.P. Morgan Healthcare Conference

- Updated clinical data from Phase 2 basket studies evaluating setmelanotide in Bardet-Biedl Syndrome (BBS) show continued weight loss for additional two patients at longer-term follow-up —
- Pivotal Phase 3 clinical trials in pro-opiomelanocortin (POMC) and leptin receptor (LEPR) deficiency obesity fully enrolled and on track for initial data anticipated in third quarter of 2019 followed by New Drug Application (NDA) filings -
- Combined pivotal Phase 3 clinical trial in BBS and Alström Syndrome on target and expected to complete enrollment in second half of 2019 -

Boston, MA — January 4, 2019 - Rhythm Pharmaceuticals, Inc. (NASDAQ:RYTM), a biopharmaceutical company focused on the development and commercialization of therapeutics for the treatment of rare genetic disorders of obesity, today announced that the company will outline expected 2019 milestones and review updated clinical data from its Phase 2 basket studies of setmelanotide in patients with BBS in a presentation at the 37th Annual J.P. Morgan Healthcare Conference on Thursday, January 10, 2019. Setmelanotide is a first-in-class melanocortin-4 receptor (MC4R) agonist.

“We expect 2019 to be an important year for Rhythm, marked by key advancements across our clinical development program for setmelanotide and culminating in preparations for the submission of our first NDA for POMC and LEPR deficiency obesity,” said Keith Gottesdiener, M.D., Chief Executive Officer of Rhythm. “Our significant achievements in 2018, including completing pivotal enrollment in our Phase 3 trials in POMC and LEPR deficiency obesity, initiating an additional pivotal trial in BBS and Alström Syndrome, and beginning to build an integrated patient community, have brought us closer to providing a first-in-class therapeutic option for people living with rare genetic disorders of obesity who have no approved treatments for reducing body weight and hunger.”

Updated BBS Data from Phase 2 Basket Studies and Pivotal Phase 3 Trial Design

Today, Rhythm announced updated clinical data from two adolescent patients in its Phase 2 basket studies evaluating setmelanotide for the treatment of BBS who previously had only short-term results (18 and 15 weeks of total treatment). Data is now available for 47 and 41 weeks of treatment. These two patients have lost 11.2% (-13.7 kg) and 15.5% (-13.7 kg) of their body weight and experienced hunger score reductions of 66% and 21%, respectively. Treatment with setmelanotide continues to be well tolerated and safety data were consistent with previous clinical studies. In total, six out of nine BBS patients enrolled in the Phase 2 basket studies have now achieved a clinically meaningful weight loss of 10% change from baseline, which is the primary endpoint for the company’s pivotal Phase 3 clinical trial, and, as previously reported, one additional patient with Type-1 diabetes responded with marked improvements in hunger score and blood sugar levels.

In December 2018, Rhythm began treating patients in a combined pivotal Phase 3 clinical trial evaluating setmelanotide in BBS and Alström Syndrome and expects to complete pivotal enrollment of at least 20 patients with BBS and at least six patients with Alström Syndrome in the second half of 2019.

Progress in Phase 3 Clinical Trials in POMC and LEPR Deficiency Obesity

Rhythm’s ongoing, separate pivotal Phase 3 clinical trials evaluating setmelanotide in POMC and LEPR deficiency obesity continue to progress, with topline data expected in the third quarter of 2019. Pending positive results, the

company expects to submit an NDA filing for each of POMC and LEPR deficiency obesity to the U.S. Food and Drug Administration (FDA) in late 2019 or early 2020.

“In 2019, Rhythm expects to make significant strides toward our vision of becoming a fully-integrated biotechnology company, as we approach the anticipated submissions of concurrent NDAs for setmelanotide in POMC and LEPR deficiency obesity,” said Murray Stewart, M.D., Chief Medical Officer of Rhythm. “As we advance toward this milestone, we are working hard to build an integrated community of patients, physicians, payors and caregivers, in order to better understand the burden of these disorders and ensure that people living with these conditions are diagnosed and readily able to access treatment.”

Additional Development Pipeline Updates

Rhythm continues to evaluate setmelanotide for the treatment of additional rare genetic disorders of obesity in its Phase 2 basket studies, including POMC and other MC4R pathway deficiency heterozygous obesities, as well as POMC epigenetic disorders. The company expects to announce updated interim data from these disorders in the first quarter of 2019. In addition, Rhythm is continuing to expand the evaluation of setmelanotide into additional MC4R pathway disorders in its Phase 2 basket studies.

Webcast Information

Rhythm will webcast its corporate presentation from the 37th Annual J.P. Morgan Healthcare Conference in San Francisco on Thursday, January 10, 2019 at 10:00 a.m. PST (1:00 p.m. EST). Live webcasts of the presentation and subsequent breakout session can be accessed under “Events & Presentations” in the Investors & Media section of the company’s website at www.rhythmtx.com. A replay of the webcasts will be available on the Rhythm website for 30 days.

About Setmelanotide

Setmelanotide is a potent, first-in-class, MC4R agonist in development for the treatment of rare genetic disorders of obesity. Setmelanotide activates MC4R, part of the key biological pathway that independently regulates energy expenditure and appetite. Variants in genes within the MC4R pathway are associated with unrelenting hunger and severe, early-onset obesity. Rhythm is currently developing setmelanotide as a replacement therapy for patients with monogenic defects upstream of MC4R, for whom there are no effective or approved therapies. The FDA has granted Breakthrough Therapy designation to setmelanotide for the treatment of obesity associated with genetic defects upstream of the MC4 receptor in the leptin-melanocortin pathway, which includes POMC deficiency obesity, LEPR deficiency obesity, BBS and Alström Syndrome. The European Medicines Agency has also granted PRiority MEDicines (PRIME) designation for setmelanotide for the treatment of obesity and the control of hunger associated with deficiency disorders of the MC4R pathway.

About Rhythm

Rhythm is a biopharmaceutical company focused on the development and commercialization of therapies for the treatment of rare genetic disorders of obesity. Rhythm is currently evaluating the efficacy and safety of setmelanotide, the company’s first-in-class MC4R agonist, in Phase 3 studies in patients with POMC deficiency obesity, LEPR deficiency obesity, BBS, and Alström Syndrome. Rhythm is dedicated to improving the understanding of severe obesity that results from specific genetic disorders. For healthcare professionals, visit www.UncommonObesity.com for more information. For patients and caregivers, visit www.LEADforRareObesity.com for more information. The company is based in Boston, MA.

Forward-Looking Statements

This press release contains certain statements that are forward-looking within the meaning of Section 27A of the Securities Act of 1933, as amended, and Section 21E of the Securities Exchange Act of 1934, as amended, and that involve risks and uncertainties, including statements regarding Rhythm's expectations regarding its expectations for 2019, anticipated timing for enrollment and design of clinical trials, the timing for filing of an NDA, and the release of results of clinical trials. Statements using word such as "expect", "anticipate", "believe", "may", "will" and similar terms are also forward looking statements. Such statements are subject to numerous risks and uncertainties, including but not limited to, our ability to enroll patients in clinical trials, the outcome of clinical trials, the impact of competition, the ability to achieve or obtain necessary regulatory approvals, risks associated with data analysis and reporting, and expenses, and other risks as may be detailed from time to time in our Annual Reports on Form 10-K and quarterly reports on Form 10-Q and other reports we file with the Securities and Exchange Commission. Except as required by law, we undertake no obligations to make any revisions to the forward-looking statements contained in this release or to update them to reflect events or circumstances occurring after the date of this release, whether as a result of new information, future developments or otherwise.

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Rhythm Pharmaceuticals:
Targeting upstream MC4R pathway defects to
transform the care of genetic obesity

Company Presentation - January 2019

Forward Looking Statements

This presentation contains certain statements that are forward-looking within the meaning of Section 27A of the Securities Act of 1933, as amended, and Section 21E of the Securities Exchange Act of 1934, as amended, and that involve risks and uncertainties, including statements regarding Rhythm's expectations for 2019 and 2020, anticipated timing for enrollment, design and completion of clinical trials, the timing for filing of an NDA, the release of results of clinical trials, expectations regarding the use of cash, and Rhythm's strategy, prospects and plans. Statements using word such as "expect", "anticipate", "believe", "may", "will" and similar terms are also forward looking statements. Such statements are subject to numerous risks and uncertainties, including but not limited to, our ability to enroll patients in clinical trials, the outcome of clinical trials, the impact of competition, the ability to achieve or obtain necessary regulatory approvals, risks associated with data analysis and reporting, our expenses, and other risks as may be detailed from time to time in our Annual Reports on Form 10-K and quarterly reports on Form 10-Q and other reports we file with the Securities and Exchange Commission. Except as required by law, we undertake no obligations to make any revisions to the forward-looking statements contained in this presentation or to update them to reflect events or circumstances occurring after the date of this presentation, whether as a result of new information, future developments or otherwise.

Transforming the Care of Patients with Genetic Obesity

The MC4R pathway is the key pathway regulating weight and hunger. Genetic defects in the MC4R pathway cause insatiable hunger, leading to severe and early-onset obesity

There are currently no effective or approved treatments for people living with rare MC4R pathway disorders

Potential First-in-Class Therapy

Setmelanotide is an MC4R agonist designed to rescue the pathway from genetic defects that occur upstream of MC4R receptor:

- **Dramatic reductions in weight and hunger observed in four rare MC4R pathway disorders:**
 - Awarded *FDA Breakthrough Therapy* and *EMA PRIME* designations
 - *Pivotal Phase 3 trials ongoing* in all four indications
- **Phase 2 development ongoing in two additional MC4R-pathway disorders:**
 - Preliminary efficacy demonstrated in heterozygous and epigenetic disorders of MC4R pathway

Multiple Avenues to Accelerate Growth

- **GO-ID Genotyping Study and TEMPO Registry:** Enable clinical study enrollment, patient identification and exploration of new genetic variants tied to MC4R loss of function
- **Rhythm Basket Study:** Facilitates rapid enrollment and exploration of setmelanotide's potential in patients with new genetic targets or syndromes tied to the MC4R pathway
- **Pipeline Expansion:** Leverage Rhythm expertise to treat additional rare genetic obesity disorders

Genetic Disorders of Obesity Impact Every Aspect of Daily Life

Meet Katy: Living with an MC4R Pathway Disorder

"It causes extreme unrelenting hunger and excessive eating. As a child...the fridge and food was controlled massively...but nobody could understand that I was desperately hungry and just wanted to stop that feeling."

3 YEARS



11 YEARS, 231 POUNDS



23 YEARS, 450 POUNDS



INFANCY:

"Normal" weight at birth, but begins to rapidly gain weight at 9 weeks

4 YEARS:

Diagnosed with POMC Heterozygous Deficiency Obesity

CHILDHOOD:

Self-isolation and missed school days
Asthmatic, increased pain and pressure on her knees make play and PE difficult

ADOLESCENCE:

Put on anti-depressants
Numbness and agonizing back pain
Abnormal pubertal development

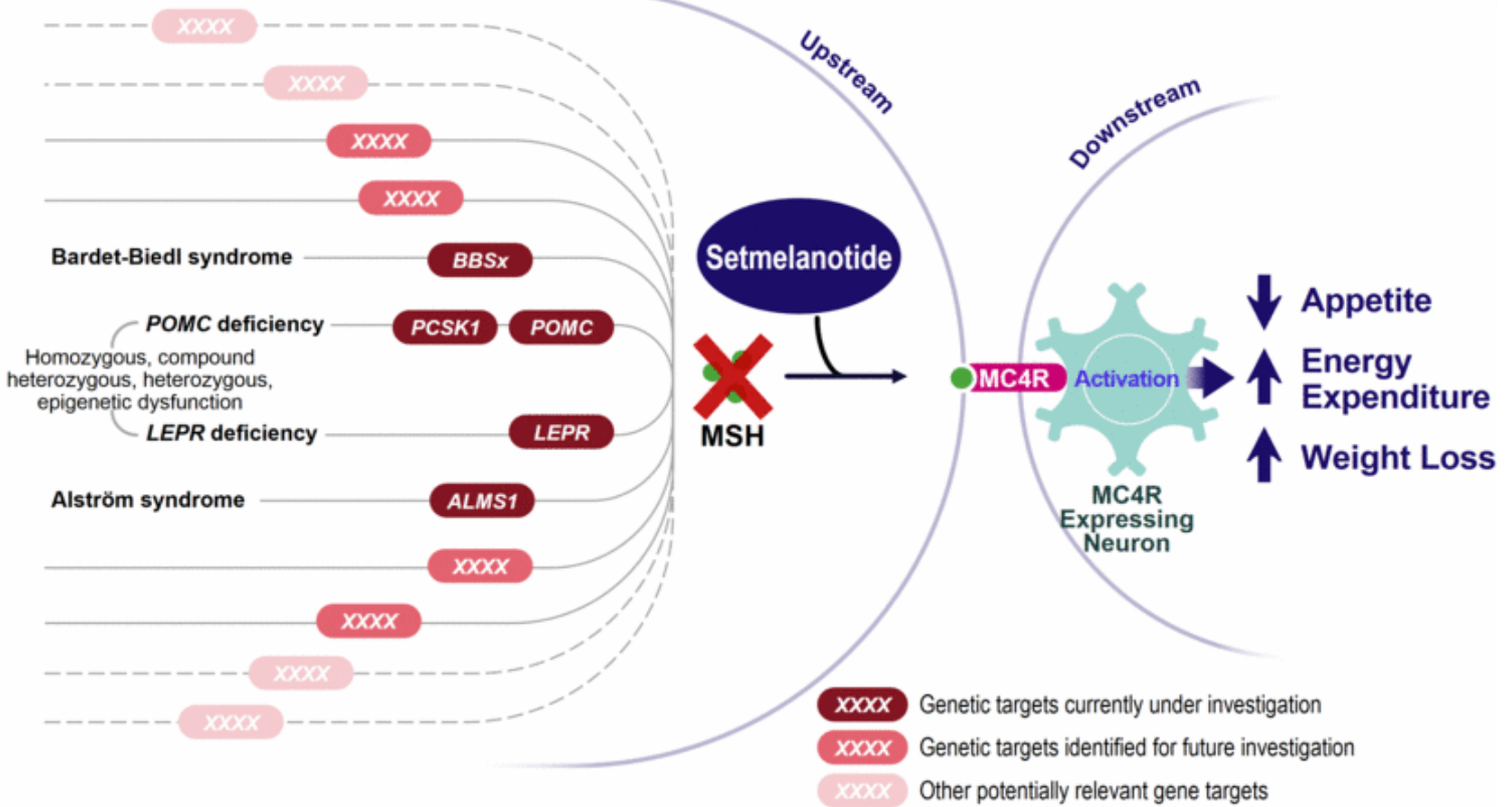
23 YEARS (CURRENT):

Sleep apnea; some cardiac issues; insulin resistance. Cracked and bleeding skin

Genetic Defects in the MC4R Pathway Cause Insatiable Hunger and Obesity

Setmelanotide Rescues the Impaired MC4R Pathway

MC4R pathway genes





**Setmelanotide: Potential First-in-
Class Replacement Therapy for
MC4R Pathway Disorders**

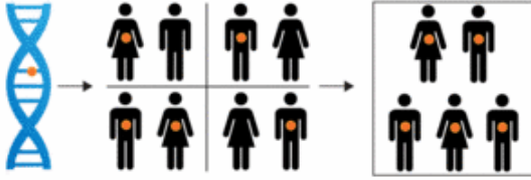
Potent first-in-class MC4R agonist



- Setmelanotide: 8 amino acid peptide with high potency (EC_{50} 0.27nM)
 - Administered by once daily subcutaneous injection
 - Retains the specificity and functionality of the naturally-occurring hormone
- Once-weekly formulation in clinical development
 - Developed in partnership with Camurus
 - Terminal half-life ~123 hours
- Previous clinical trials with approximately 300 obese patients demonstrated statistically significant weight loss and favorable tolerability profile
- IP: Composition of matter in all major markets
 - US patent, with possible Hatch-Waxman extension, to 2032
- Highly competitive cost of goods sold

* NOAEL: No-observed-effect-level

Phase 2 Basket Study



Designed to facilitate rapid progress to proof of concept (POC) in potential new indications for setmelanotide



3 month proof-of-concept study phase:
Weight loss, hunger and other metabolic parameters are measured



If significant weight loss and acceptable safety and tolerability is demonstrated

1 year open-label, single-arm, proof of concept extension studies

Indications enrolled in Phase 2:

- POC Achieved, Phase 3 initiated: POMC, LEPR, BBS & Alström Syndrome
- Pre-POC, Phase 2 ongoing: MC4R pathway heterozygous obesity, POMC epigenetic obesity
- New Indications: Additional genetic targets upstream of MC4 receptor

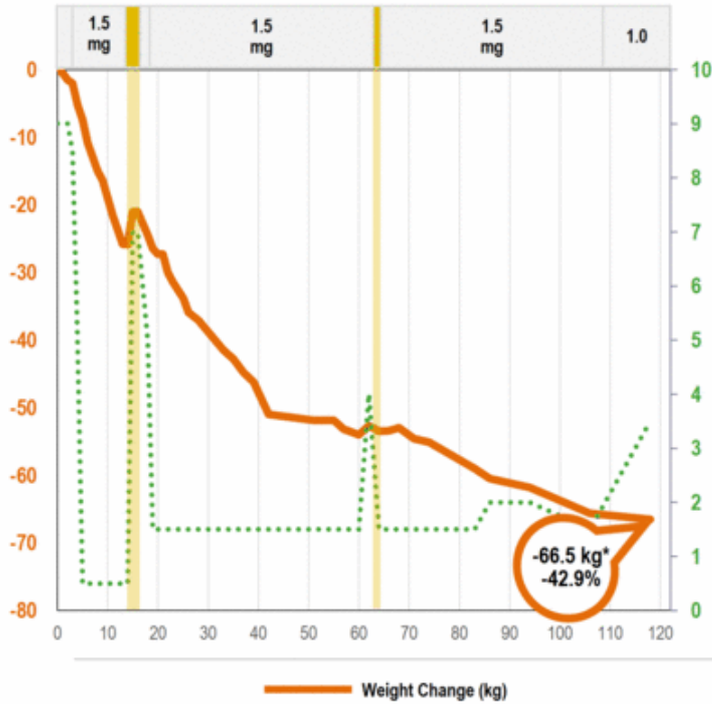
Close coordination with GO-ID Genotyping Study:

- Patients with genetic mutations identified from GO-ID study eligible to enroll in basket study

POMC and LEPR Phase 2 Studies: Patients #1

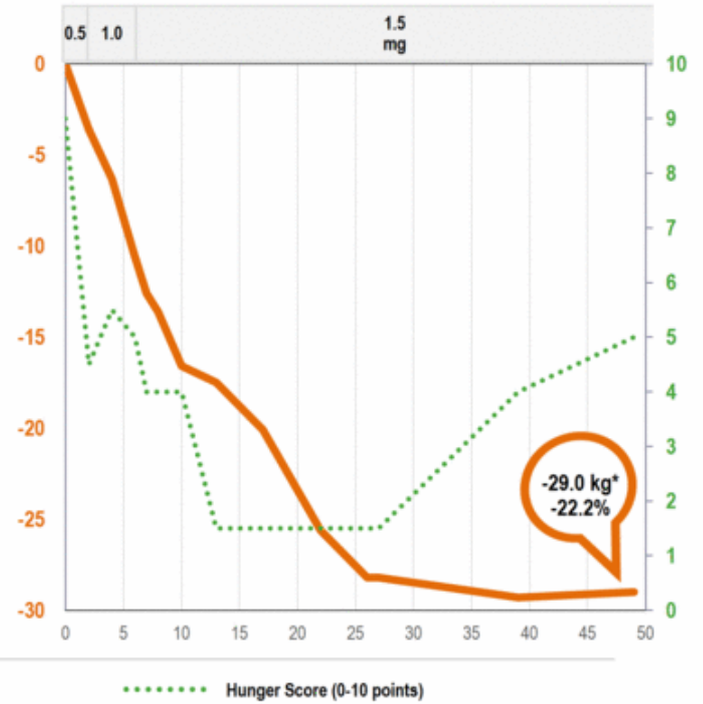
POMC Patient #1⁽¹⁾

20 yr old female
 Starting Weight = 155.0 kg
 Starting BMI = 49.8 kg/m²



LEPR Patient #1⁽²⁾

23 yr old male – LEPR Mutation
 Starting Weight = 130.6 kg
 Starting BMI = 39.9 kg/m²



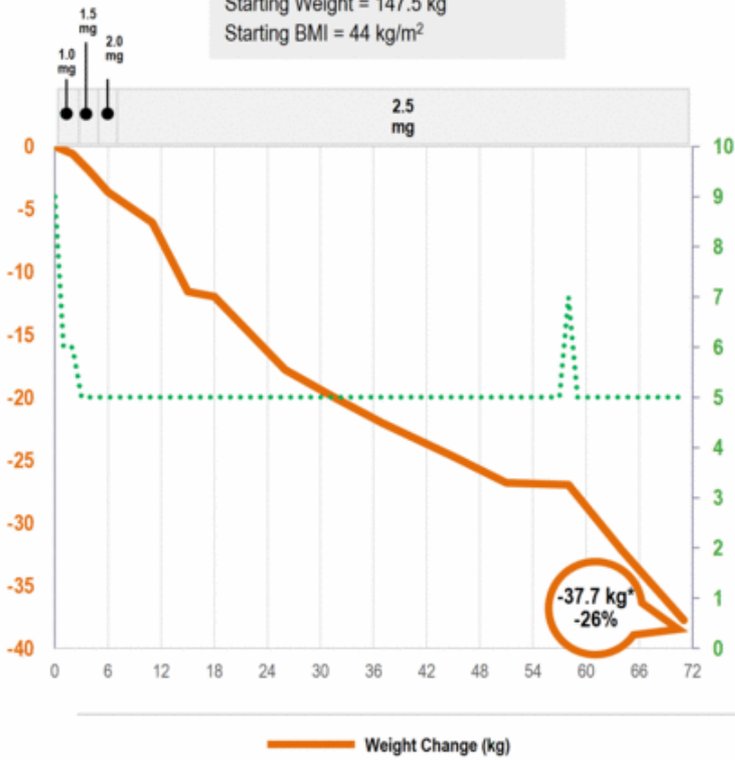
* Figures represent cumulative weight lost in kgs | Not all patients had similar responses

(1) Kühnen, et. al, Proopiomelanocortin Deficiency Treated with a Melanocortin-4 Receptor Agonist. *N Engl J Med.* July 2016 . Yellow vertical bars represent intervals with dose withdrawal or modifications
 (2) Biebermann, et al. MC4R Agonism Promotes Durable Weight Loss in Patients with Leptin Receptor Deficiency. *Nat Med.* 2018 May 7.

BBS and Alström Syndrome Phase 2 Studies: Patients #1

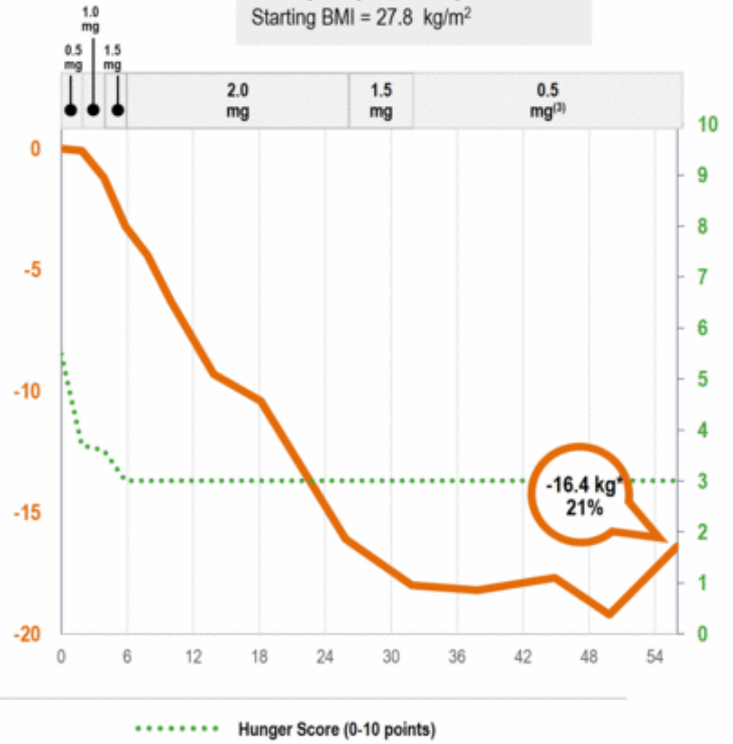
BBS Patient #1⁽¹⁾

25 yr old male - BBS1 Mutation
 Starting Weight = 147.5 kg
 Starting BMI = 44 kg/m²



AS Patient #1⁽²⁾

12 yr old male
 Starting Weight = 78.6 kg
 Starting BMI = 27.8 kg/m²



* Figures represent cumulative weight lost in kgs | Not all patients had similar responses

- (1) Haws et al. Effect of the Melanocortin-4 Receptor Agonist Setmelanotide on Obesity and Hyperphagia in Individuals Affected by Bardet-Biedl Syndrome. ESPE 2018, Sept. 28, 2018.
- (2) Han et al. Effect of Setmelanotide (MC4R Agonist) on Obesity and Hunger in Individuals With Alström Syndrome. ObesityWeek 2018, November 15, 2018.
- (3) From week 49 onwards, the dose was lowered to 0.5 mg five times a week

Phase 2 BBS and Alström Results¹ Support Advancement to Phase 3

	Variant	Sex	Age, years	Treatment, weeks	Weight Change	Hunger Score Reduction
Patient 1	BBS1	Male	25	65	-26% (-37.7 kg)	44%
Patient 2	BBS2	Female	61	55	-15% (-14.5 kg)	86%
Patient 3	BBS10	Female	16	64	-24% (-29.1 kg)	N/A
Patient 4	BBS12	Female	17	50	-24% (-23.1 kg)	83%
Patient 6	BBS5	Female	16	47	-11.2% (-13.7 kg)	66%
Patient 7	BBS4	Female	14	41	-15.5% (-13.7 kg)	21%

- **6 of 9 BBS patients achieved clinically significant weight loss of >10% change from baseline**
 - Patients 6 and 7: Updated data (Jan. 2019) demonstrate longer-term weight loss and hunger score reductions
- **3 patients discontinued treatment**
 - Patient 5 (pediatric patient with BBS1 variant and type 1 diabetes) experienced 53.3% reduction in hunger and reduction in hemoglobin A1c (10.1% to 7.6%) before withdrawing
 - Two patients (one non-genetically confirmed) withdrew due to lack of weight loss

(1) Haws RM et al. Effect of the Melanocortin-4 Receptor Agonist Setmelanotide on Obesity and Hyperphagia in Individuals Affected by Bardet-Biedl Syndrome. ESPE 2018. Abstract RFC6.3. Updated data for pts. 6 & 7 announced by Rhythm in January 2019.

Heterozygous and Epigenetic Obesity Disorders

- Focus on most debilitated patients
 - **MC4R pathway heterozygous deficiency**: one loss of function allele
 - **POMC epigenetic disorders**: disease caused by DNA modifications that can change gene expression without altering DNA
- Initial preliminary data as of June 2018:
 - **MC4R pathway heterozygous deficiency**: positive initial weight and hunger responses in two evaluable patients
 - **POMC epigenetic disorders**: positive initial weight and hunger response in one evaluable epigenetic patient
- Updated interim data expected in first quarter of 2019
- Plan to continue enrolling additional patients, in order to better identify those likely to benefit from setmelanotide and inform next steps

Safety and Tolerability

Setmelanotide has been evaluated in over 300 obese subjects

- Setmelanotide has been generally well-tolerated
- Most AEs are mild and non-mechanism based:
 - Mild injection site reactions
 - Darkening of skin (tanning) and skin lesions, mediated by the closely related MC1 receptor (the natural “tanning” receptor)
- Discontinuations are rare; no increase in CV parameters

All pivotal trials share a common design:

- Patients aged ≥ 6
- ~1-year trial duration, with long-term extension
- Dosage: Daily subcutaneous injection
- 1^o endpoint: categorical analysis of responders⁽¹⁾
- Key 2^o endpoints include mean weight loss and hunger measures
- Supplemental patients enrolling beyond pivotal cohorts to generate additional data

POMC and LEPR Phase 3 Trials

- Pivotal cohorts (n~10) fully enrolled for each trial
- Top-line data from both trials expected in 3Q19

BBS/Alström Combined Phase 3 Trial

- Pivotal cohort (n~30), with at least 20 BBS patients and 6 Alström patients
- 14-week placebo-controlled period, followed by completion of active open-label treatment for total duration of ~52 weeks
- Completion of pivotal enrollment expected in 2H19

⁽¹⁾ Percentage of patients who have at least a 10% reduction in body weight at approximately one year.



**Building a Community for the
Treatment of Rare Genetic Disorders
of Obesity**

Epidemiology Suggests Significant Market Opportunity

POMC & LEPR Deficiency Obesity: Non-Syndromic Disorders



Patients diagnosed after genetic screening.

BBS and Alström Syndrome: Syndromic Disorders



Patients often known to the medical system.
E.g. Over 450 BBS patients enrolled in CRIBBS registry

CLINICAL EPIDEMIOLOGY

Up to 5,000 patients
in U.S.⁽¹⁾

POMC: 100-500*

LEPR: 500-2,000*

BBS: 1,500-2,500*

Alström: 500-1,000 (worldwide)

GENETIC EPIDEMIOLOGY⁽²⁾

~13,000 individuals in the U.S. with POMC or LEPR -- a **five-fold increase** over the clinical epidemiology estimate.

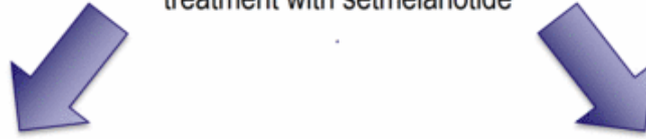
* The patient numbers above are based on company estimates

(1) Rhythm believes that the addressable patient population in Europe is at least as large as in the U.S.

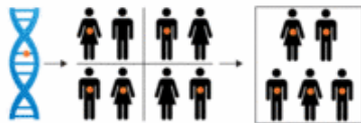
(2) Ayers et al. [Melanocortin-4 Receptor Pathway Dysfunction in Obesity: Patient Stratification Aimed at MC4R Agonist Treatment](#). *J Clin Endocrinol Metab.* 2018 May 2. doi: 10.1210/je.2018-00258. JCEM citation



Identifies individuals with rare genetic disorders of obesity who may be eligible for further study and treatment with setmelanotide



Rhythm-Sponsored Phase 2 Basket Study



Designed to facilitate rapid progress to proof of concept in potential new indications for setmelanotide



TRACING THE EFFECT OF THE MC4R PATHWAY IN OBESITY

Commitment to understanding the ongoing impact and burden of disease on patients and their caregivers



Identifies individuals with rare genetic disorders of obesity who may be eligible for further study and treatment with setmelanotide

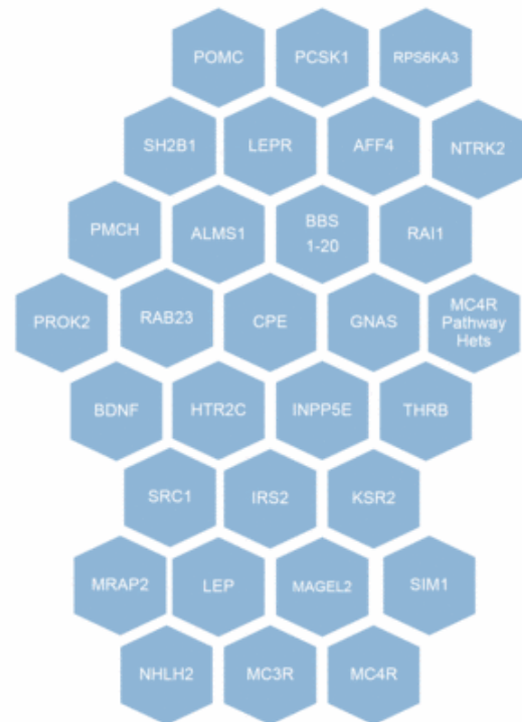
- Genotyping study of individuals with obesity ≥ 2 years of age
- Recruitment via outreach to HCPs caring for patients with severe obesity
- Expected enrollment of $\geq 9,000$ patients from more than 140 centers worldwide
- Genotyping panel includes over 100 genes associated with genetic obesity
- Individuals identified with mutations in MC4R pathway genes may be enroll in future clinical trials, observational trials, or TEMPO registry
- Goal: Increase the frequency of genetic testing and shift practice patterns among treating physicians



TRACING THE EFFECT OF THE MC4R PATHWAY IN OBESITY

- Commitment to understanding the ongoing impact and burden of disease on patients and their caregivers
 - Facilitates enhanced understanding of these conditions in the medical community
 - Builds upon ongoing collaborations with existing patient registries (e.g., CRIBBS for BBS)
- Target enrollment of ~1,000 patients
- Genetic screening through GO-ID
- Potential enrollment in Basket study

TEMPO is for individuals with specific variants in the MC4R pathway genes, that include at least one of the following:



Rhythm Achieved Key Milestones in 2018



- ✓ Completed enrollment in POMC & LEPR Phase 3 trials
- ✓ Announced updated positive Phase 2 BBS & Alström results
- ✓ Breakthrough Therapy and EMA Prime designation expanded to include BBS & Alström
- ✓ Enrolled first patient in pivotal Phase 3 study in BBS & Alström
- ✓ Advanced Phase 2 trials of MC4R pathway heterozygous deficiency and epigenetic disorders
- ✓ Launched TEMPO registry to gather information about people affected by rare genetic disorders of obesity
- ✓ In-licensed RM-853, potential therapy for Prader-Willi
- ✓ Completed follow-on offering, extending expected cash runway to second-half of 2020

Rhythm Expects Significant Growth in 2019 and 2020



- 1Q19** Updated interim data for MC4R pathway heterozygous and epigenetic obesity disorders
- 3Q19** Topline data from both POMC and LEPR Phase 3 studies
- 4Q19-1Q20** Initial NDA submission filings for setmelanotide in POMC and LEPR
- 2H19** Complete pivotal enrollment in BBS and Alström Phase 3 study
- 2019** Update on ongoing efforts to increase patient identification
- 2019** Expand Phase 2 basket studies into additional MC4R pathway disorders
- 2020** Topline data from BBS and Alström Phase 3 study

Living with the Insatiable Hunger and Obesity that Characterize BBS



“As our daughter Lucy grows older, the weight obstacle of BBS looms more ominously before us, hindering much of what we do. Lucy measures most of her activities and much of her happiness around when her next meal is, what food is available where, and it is an overwhelming task to help her take control. Weight affects so much in her life and we are working so hard to fight to give her a more fulfilling life without the confines that obesity presents.”

-Shawni, mother to Lucy, a child living with BBS





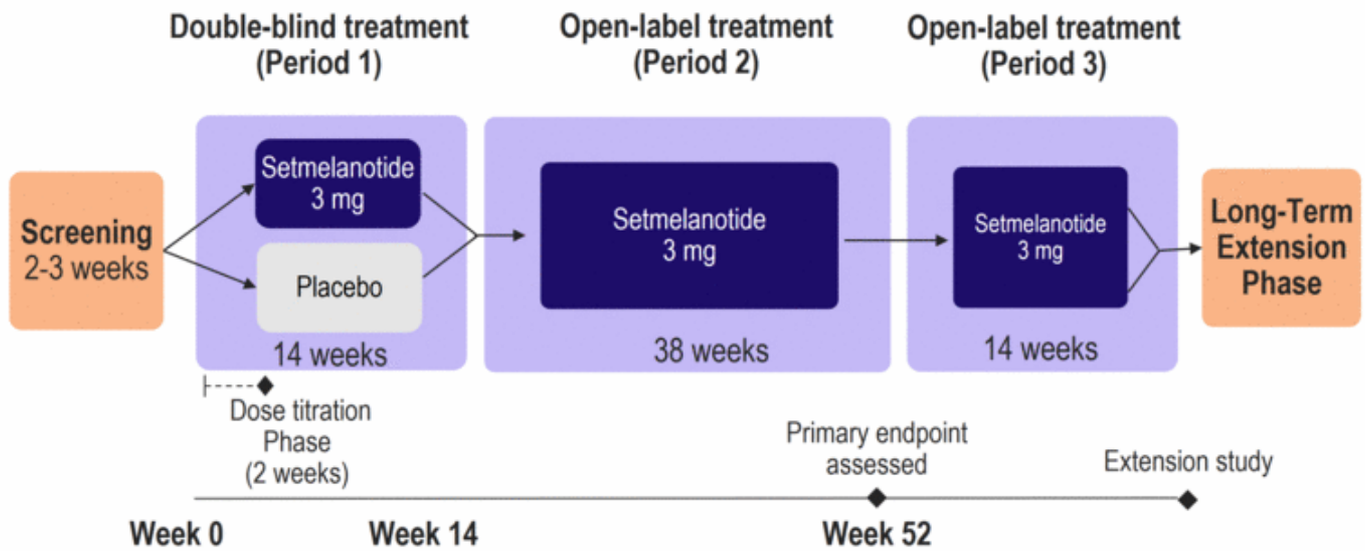
Appendix

Rhythm Pipeline

Indication	PRE-CLINICAL	PHASE 1	PHASE 2	PHASE 3
SETMELANOTIDE				
POMC Deficiency Obesity	▶			
LEPR Deficiency Obesity	▶			
Bardet-Biedl Syndrome	▶			
Alström Syndrome	▶			
POMC & Other MC4R Pathway Heterozygous Deficiency Obesities	▶			
POMC Epigenetic Disorders	▶			
RM-853 (GHRELIN O-ACYLTRANSFERASE INHIBITOR)				
Prader-Willi Syndrome	▶			

Rhythm is currently assessing opportunities to further evaluate setmelanotide in PWS and plans to pursue these in parallel with the development of RM-853.

Bardet Biedl & Alström Syndrome: Phase 3 Study Design

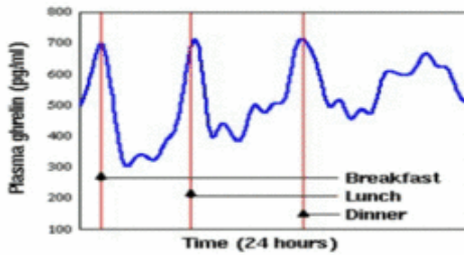


Primary Endpoint: Percentage of patients (12 years of age and older at Week 0) who have at least a 10% reduction in body weight at Week 52

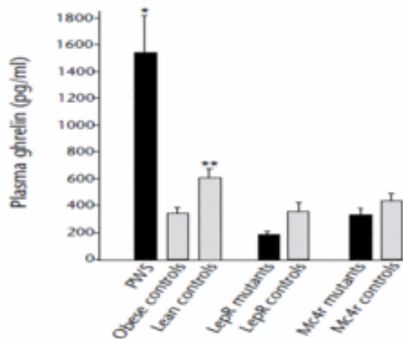
Protocol	<ul style="list-style-type: none">▪ Final comments from FDA have been incorporated
Study enrollment milestones	<ul style="list-style-type: none">▪ <u>First patient with BBS enrolled</u>
Study enrollment goals	<ul style="list-style-type: none">▪ Enrollment <u>minimum</u>:<ul style="list-style-type: none">• 20 BBS patients• 6 Alström Syndrome patients
Centers of excellence	<ul style="list-style-type: none">▪ Marshfield Clinic Health System<ul style="list-style-type: none">• Bob Haws, MD▪ University College London Great Ormand Street Institute of Child Health<ul style="list-style-type: none">• Phil Beales, MD
Confirmed Countries	<ul style="list-style-type: none">▪ United States, United Kingdom, Canada, Netherlands, Spain

RM-853: Potent, Orally Available GOAT Inhibitor for PWS

Ghrelin is tightly correlated with hunger signals throughout the day



People with PWS have higher ghrelin levels



- GOAT is key enzyme involved in producing active ghrelin
- Blocking GOAT results in:
 - Lower levels of active ghrelin, *and*
 - Increased levels of DAG, a ghrelin precursor believed to have independent beneficial effects
- In preclinical studies with high fat-fed mice, RM-853 prevented body weight gain and reduced fat mass, with favorable PK, PD and safety profile
- Development plan complements and expands ongoing efforts
 - RM-853 IND filing expected Q1 2020
 - Still planning to evaluate setmelanotide in PWS, in parallel with development of RM-853
 - Future Phase 2 trial of setmelanotide may evaluate longer durations of treatment, higher doses, younger patients and operational limitations of prior trial
 - Given setmelanotide and RM-853's distinct mechanisms of action, will explore opportunities to evaluate in combination, as there may be complementary effects

Cash expected to be sufficient to Fund Operations into 2H '20

SHARES OUTSTANDING
as of 9/30/2018

34,382,525
(basic)
36,857,315 million
(fully-diluted)

**CASH, CASH EQUIVALENTS AND
SHORT TERM INVESTMENTS**
as of 9/30/2018

\$ 272.4 million

Raised \$163M in net proceeds in June 2018 follow-on offering




rhythm