

Zacks Small-Cap Research

Sponsored – Impartial - Comprehensive

Brian Marckx, CFA
bmarckx@zacks.com
Ph (312) 265-9474

scr.zacks.com

10 S. Riverside Plaza, Chicago, IL 60606

Biomerica Inc

(BMRA-NASDAQ)

BMRA: Legacy Biz Slows in Q4. Operational Highlights Include InFoods Enrolling, Ex-Walgreens CMO Joins SAB (and we're intrigued)

We use sum of the parts to value BMRA. Comparable P/S and P/B values the base business at ~\$3.00/share. Applying 11x to our \$8.4M 2024 forecasted InFoods revenue and discounting back to the present at 15%/year, results in InFoods present value of approximately \$35M, or \$4.00/share. BMRA TOTAL VALUE = \$7/share

Current Price (09/06/18) **\$3.80**
Valuation **\$7.00**

OUTLOOK

Biomerica reported financial results for their fiscal 2018 fourth quarter ending May 31, 2018. What looked like the front end of a recovery in the top-line through the first nine months ended with a whimper, with revenue in the final quarter of fiscal 2018 dropping 22% yoy and 18% sequentially. While our model had revenue inching up 6% for the full-year reflecting an expectation that rebounding sales in Asia would more than offset ongoing softness in both Europe and the U.S., that was far from the case. Instead, the nearly across-the-board (Latin America was the sole bright spot) weakness in Q4, which most notably included a 42% yoy contraction in Asia, resulted in annual sales falling almost 4%. Major milestone with first patient enrolled in InFoods Study #1 (endpoint determination). H. pylori studies also ongoing – could have some announcements in the near-term.

We're intrigued by the appointment of Harry Leider to BMRA's SAB. At Walgreens Mr. Leider had responsibility for overseeing the evaluation of new technologies aimed at new services. While pure speculation on our part, we wonder if the appointment to BMRA's SAB may signal that management is looking at the feasibility of the retail pharmacy channel for InFoods (think Theranos' strategy – kind of, but without the bad stuff).

SUMMARY DATA

52-Week High **\$9.32**
52-Week Low **\$2.68**
One-Year Return (%) **32.84**
Beta **-0.66**
Average Daily Volume (sh) **8,120**

Shares Outstanding (mil) **9**
Market Capitalization (\$mil) **\$32**
Short Interest Ratio (days) **N/A**
Institutional Ownership (%) **2**
Insider Ownership (%) **26**

Annual Cash Dividend **\$0.00**
Dividend Yield (%) **0.00**

5-Yr. Historical Growth Rates
Sales (%) **N/A**
Earnings Per Share (%) **N/A**
Dividend (%) **N/A**

P/E using TTM EPS **N/A**
P/E using 2019 Estimate **N/A**
P/E using 2020 Estimate **N/A**

Zacks Rank **N/A**

Risk Level

Type of Stock
Industry

Above Avg.,
Small-Growth
Med-Drugs

ZACKS ESTIMATES

Revenue

(in millions of \$)

	Q1	Q2	Q3	Q4	Year
	(Aug)	(Nov)	(Feb)	(May)	(May)
2018	1.45 A	1.61 A	1.38 A	1.13 A	5.56 A
2019	1.33 E	1.44 E	1.68 E	1.53 E	6.00 E
2020					6.39 E
2021					6.88 E

Earnings per Share

	Q1	Q2	Q3	Q4	Year
	(Aug)	(Nov)	(Feb)	(May)	(May)
2018	-\$0.02 A	-\$0.03 A	-\$0.04 A	-\$0.08 A	-\$0.17 A
2019	-\$0.06 E	-\$0.06 E	-\$0.05 E	-\$0.06 E	-\$0.22 E
2020					-\$0.37 E
2021					-\$0.26 E

Zacks Projected EPS Growth Rate - Next 5 Years % **N/A**

WHAT'S NEW.....

Q4 2018: *Legacy Biz Slows in Q4. Operational Highlights Include InFoods Enrolling, Ex-Walgreens CMO Joins SAB (and we're intrigued)...*

Biomerica reported financial results for their fiscal 2018 fourth quarter ending May 31, 2018. What looked like the front end of a recovery in the top-line through the first nine months ended with a whimper, with revenue in the final quarter of fiscal 2018 dropping 22% yoy and 18% sequentially. While our model had revenue inching up 6% for the full-year reflecting an expectation that rebounding sales in Asia would more than offset ongoing softness in both Europe and the U.S., that was far from the case. Instead, the nearly across-the-board (Latin America was the sole bright spot) weakness in Q4, which most notably included a 42% yoy contraction in Asia, resulted in annual sales falling almost 4%.

- U.S. revenue has been weak and disappointing as of late. As we noted in our update in April, fiscal Q3 has historically been a relatively strong quarter for BMRA's U.S. business – and while that again proved to be true, the \$208k in U.S. revenue generated in Q3'18 was the lowest Q3 U.S. number since at least 2010. U.S. revenue in Q4'18 was \$152k – while up 16% yoy, it's the second lowest Q4 U.S. revenue number since at least 2010. BMRA indicated that the decline in U.S. sales relates to lower purchasing (of their flagship EZ Detect colon disease test) from a certain drug store customer that, unlike some years, decided not to conduct a colorectal cancer screening program during fiscal 2018. We remain optimistic, however, that U.S. activity will come back and believe a possible catalyst to U.S. sales growth could come from the new h. pylori test candidate which just enter clinical studies
- Europe has similarly been weak and softer than what we had anticipated. We noted in our Q3 update that weakness in Europe could continue and put a damper on BMRA's full-year results – but also noted that we thought European FY'18 revenue would not contract by more than single digits. While we were prescient relative to ongoing weakness - with Europe revenue down 19% yoy and 5% sequentially in Q4, we slightly underestimated the degree as full-year revenue, down 10%, did manage to eke out a double-digit decline. Lower private label sales were cited as a contributor to the relative weakness in Europe.
- Asia rounded out the disappointment in Q4 with revenue down 42% yoy, down 37% sequentially and, at \$394k, the weakest showing from that territory since Q3'16. Fortunately, relatively robust activity through the first nine months of the year, pushing revenue up 22% YTD through Q3, help mute the Q4 weakness. Asia revenue, which accounts for ~45% of BMRA's total, was up 4% for the full year. The softness in Asia in Q4 is somewhat confounding as revenue growth had been on a fairly steep trajectory since late-fiscal 2015, which was shortly after BMRA restructured their distribution channel in China. We had also anticipated incremental revenue growth from the launch of EZ Detect, which received China FDA (CFDA) approval in January of this year (i.e. late-fiscal Q2'18).

Total revenue fell 22% yoy and 18% qoq to \$1.13M in Q4 – which is the lowest quarterly revenue since fiscal Q1'15. For the full-year, revenue was \$5.56M – while down 4% from FY2017, it is up from 2014 (\$5.12M), 2015 (\$4.96M) and 2016 (\$5.14M). We also note that revenue through the first nine months of fiscal 2018 was the highest of any prior year since 2013. Our point being that, if Q4 was an outlier and fundamentals have not materially changed, revenue growth from BMRA's current products is likely to reemerge in the near-term.

U.S. revenue, at \$152k was up 16% yoy – although that overstates the implied strength as the prior-year comp was the lowest quarterly revenue from the U.S. since at least fiscal 2010. U.S. sales have trended relatively soft since 2015 when they fell 17% to \$1.0M in that year. They then dipped another 4% in 2016, 12% in 2017 and finished 2018 at \$685k, down another 22%, and the lowest level since at least 2010 (i.e. the furthest back that we looked).

While difficult to predict, we model U.S. sales to stabilize at or around recent levels (related to the company's current product portfolio) over the near-term. A possible catalyst to U.S. sales growth could come from a new h. pylori test, clinical trials for which recently commenced enrollment (and which we discuss in more detail below).

And with the recent disclosure and announcement of initial clinical trial design and collaboration to conduct clinical trials with two major U.S. university research centers, we recently began modeling InFoods. As we discuss in more detail below, we are modeling initial contribution from InFoods in 2022. We continue to believe that eventual FDA clearance of this novel IBS product would result in a significant increase in Biomerica's U.S. sales and provide the majority of total revenue growth from that point into the foreseeable future.

Meanwhile, **Q4 revenue from Asia** was \$394k – down 42% yoy and down 37% from Q3 of this year. Q4 revenue represented just 55% of the average through the first nine months of fiscal 2018, which was up 22% from the comparable period in 2017. The about-face in Q4 is surprising – and we have no insight for the weakness. But, as

we have noted in the past, while Asia has been a territory which has historically experienced relatively high short-term sales volatility, longer-term trends have pointed towards regular revenue growth.

And, as Asia has grown to account for a larger proportion of total revenue, the recent strength from this territory has been the most significant catalyst to driving BMRA's topline. For historical context, Asia generated just \$1.0M in sales (~21% of total revenue) in 2015 - this grew 71% to \$1.7M in 2016 and another 39% to \$2.4M in 2017. In fact, Asia was effectively the only reason why total revenue posted positive growth in 2016 and is credited with almost 90% of the topline growth in 2017.

In the most recent year, Asia sales grew 22% through the first nine months and despite the cliff-dive in Q4, still managed to eke out 4% growth in FY2018. And among the territories which did post positive revenue growth, Asia contributed 86% through the first nine months and 40% for the full-year in 2018. Given the outsized contribution from Asia (which now accounts for 45% of total revenue) even incremental growth from current levels in this territory will have a meaningfully positive effect – but even incremental contraction will have the opposite effect.

As noted, we had hoped EZ Detect (over-the-counter fecal occult blood (FOB) test for colorectal cancer) would have been a positive catalyst – although it's possible we were a little too optimistic relative to timing. We continue to like the fundamentals in China as it relates to BMRA's flagship product. While Biomerica has never publicly disclosed product-specific sales numbers, we believe EZ Detect is one of the (if not the #1) best-selling products for the company. We also think Asia could be particularly receptive to the product given certain cultural principles in many parts of Asia related to hygiene which may discourage use of FOB tests which require fecal handling.

Recent initiatives aimed at increasing colorectal cancer screening – by any modality – also support fundamentals for EZ Detect. Many parts of Asia, including in China, have relatively low rates of compliance to recommended colorectal cancer screening guidelines. For context, while ~65% of Americans adhere to CRC screening guidelines, studies indicate that compliance is only ~40% in Shanghai, China. Studies have shown that one of the most effective ways to increase CRC compliance is through providing more testing options. So, for all of these reasons, we think Asia could represent a substantial growth opportunity for BMRA's EZ Detect product.

Relative to Europe – sales from this territory showed recent signs of growth, inching up 3% in 2017 following a 20% slide the prior year. While the mid-single digit growth through the first six months of 2018 was promising and an encouraging signal that European sales growth might be accelerating, that was more than offset by weakness in the second half of the year which saw sales drop more than 22%. For the full year, European revenue was down 10% to \$2.0M, the lowest level since at least 2010.

Europe accounts for about 36% of total sales, making it the company's second most important market and causing any meaningful variability to have a significant influence on overall financial performance. And while the recent indications that sales in that territory may be gaining traction were encouraging, the lack of sustained growth, which significantly affected BMRA's topline performance over the last three years, has us questioning whether there is much upside in this area.

We now model flat to slightly negative sales in Europe related to BMRA's current product portfolio and think opportunity for sustainable growth may mostly hinge on new product launches. While we have yet to model any assumed contribution from InFoods (or any other new products), that could soon change - depending on BMRA's future strategic objectives as well as if we feel there is enough information to make (comfortably) informed projections about certain commercializability-related gating factors. At this point we have no information or insight into if or when BMRA might consider targeting markets outside of the U.S. or if they do, which areas of the world they would focus on next. Europe, however, would be our best-guess as a potential front-runner if management does eventually look to expand OUS with InFoods given not only the economic similarities of most of the highly developed European countries with that of the U.S. but, perhaps more importantly, diets that are (generally) similar to that of most Americans.

Gross Margin, Operating Expenses

Gross margin averaged a relatively healthy 33.6% through the first nine months of 2018. Gross margin was 23.7% and 31.5% in Q4 and FY'18, compared to 30.3% and 34.9% in the prior year periods. GM is negatively affected by lower sales volumes – as less fixed costs are absorbed.

OpEx was \$912k, or 81% of revenue, in Q4 and \$3.2M, or 58% of total revenue in FY2018 – which compares to \$791k (55%) and \$3.0M (51%) in the prior year. SG&A was flat through 2018 as compared to the prior year despite expanded activity in both operational and product development areas. We do, however, continue to model opex to

increase at a higher rate than that of revenue growth through at least fiscal 2019 (ending May 2019) as a result of increasing development activity related to InFoods, and to a lesser extent, the h.pylori program.

Cash

BMRA raised \$1.3M, net, during fiscal 2018 including \$981k in Q4, from the sale of common shares via their ATM program. Another \$171k (net) was raised subsequent to year-end. Cash balance at the close of FY2018 was \$1.2M, up from \$597k at the end of Q3'18. Cash used in operating activities in the three and twelve months ending May 31, 2018 was \$322k and \$1.2M (\$568k and \$1.2M, ex-changes in working capital), respectively. Another \$56k and \$130k was used in investing activities in the same periods.

In June 2017 Biomerica filed an S-3 registration statement with the SEC (which became effective July 20th), registering for sale (up to) \$45M in common stock. Then on December 1st the company entered into an ATM agreement with B. Riley FBR, authorizing the sale of up to \$7M of common shares. Through FY2018 342k shares have been sold under shelf, representing net proceeds of ~\$1.28M. As we noted following filing of the registration statement, we think proceeds would almost certainly be mostly targeted towards advancement and further development, including clinical validation, of InFoods.

Additional, non-dilutive funds, could come from BMRA's agreement with Telcon Pharmaceuticals (fka Celtis Pharm Co.) of S. Korea which calls for that company to pay Biomerica up to \$1.25M in exclusivity fees based on "certain milestones including Biomerica's starting clinical trials in the United States, receipt of US FDA clearance and Celtis' first sales of IBS Products in Korea". The agreement was initially cancellable if BMRA had not obtained FDA clearance/approval of InFoods by December 31, 2017 but that deadline was subsequently extended until December 31, 2019.

Operational Update (see Appendix for background info)

Harry Leider (ex-Walgreens CMO) Joins BMRA's SAB

While most of the operational-related excitement relates to InFoods and h.pylori product development – and which we talk about below – we think the recent appointment of Harry Leider deserves mention. Mr. Leider, who served as Chief Medical Officer of Walgreens from November 2015 to May 2018, joined BMRA's Strategic Advisory Board in April. Notably, increasing competition in the retail pharmacy space has created some significant changes in this market. Along with greater M&A activity, retail pharmacies have branched out their portfolio of products and services as they look for new ways to grow revenue. This includes expanding patient care services – such as providing wellness, screenings, vaccinations and treating minor injuries. At Walgreens Mr. Leider had responsibility for overseeing the evaluation of new technologies aimed at new services. While pure speculation on our part, we wonder if Mr. Leider's appointment to BMRA's SAB may signal that management is looking at the feasibility of the retail pharmacy channel for InFoods (think Theranos' strategy – kind of, but without the bad stuff).

InFoods Study 1 Commences Enrollment

In late-June 2018 BMRA announced that the first patient had enrolled in the InFoods Endpoint Determination study.

As a reminder, two clinical studies will be conducted; the purpose of the first study, which will include approximately 180 subjects and with an expected duration of 9 – 14 months, is to identify the primary endpoint to be used in the second study. It is anticipated that the second study will serve as the pivotal study and primary support for an eventual 510(k) filing.

Study 1, titled simply (as of now, anyway) "Endpoint Determination Study Protocol" was first posted on clinicaltrials.gov on March 9, 2018 (NCT03459482, link: <http://bit.ly/2H9RTTP>).

Particular points of interest related to Study 1 as detailed in the posting include:

- RCT-design and will incorporate triple blinding (i.e. patient, provider and investigator)
- Targeting enrollment in each arm (i.e. treatment vs. sham) of 90 patients and a minimum of 30 patients in each IBS classification of: IBS-D (diarrhea), IBS-C (constipation) and IBS-M (mixed)
- Arms
 - o Treatment (i.e. food elimination diet): patients will be given an elimination diet based upon foods with a positive antibody profile in the InFoods IBS test. The elimination diet will also exclude any and all foods to which the subject has a known IgE allergy and foods the subject already currently eliminates

- Sham: patients will be given a "Sham" elimination diet. The sham diet will eliminate the same number of foods but none of the actual foods to which the patient had a positive antibody profile in the InFoods IBS test. The sham diet will also eliminate any and all foods to which the subject has a known IgE allergy and foods the subject already currently eliminates
- Inclusion criteria includes
 - Meets Rome III or Rome IV IBS diagnostic criteria
 - Respond "No" to IBS Adequate Relief (IBS-AR) in the past week at the screening visit #1
 - Score between ≥ 3 and <7.5 on the Abdominal Pain Intensity Assessment (IBS_API) based on a weekly average of worst daily (in past 24 hours) abdominal pain on a 0 to 10 point scale
 - A positive IgG antibody response for at least one food in the InFoods IBS panel
 - Patients who are on stable (> 3 months) doses of medications or treatments for their IBS (e.g., probiotics, fiber, Viberzi, Linzess, Amitiza, Alosetron, Plecanatide, anticholinergics, antidepressants, Zofran bile acid sequestrants, or anti-diarrheals) will be allowed to continue their medications as long as no change in treatment is planned for the duration of the study and no dose adjustment is made during the duration of the study
- Exclusion criteria includes
 - Patients who have used Rifaximin in the past 3 months
 - Patients engaged in another type of diet therapy i.e. FODMAP
 - Chronic pain from other conditions besides IBS

Our Comments on Study 1 Design:

Is of robust design, RCT and triple blinding. Clearly Study 1 was designed to serve as a template for a pivotal study. Inclusion and exclusion criteria all appear to be consistent with designs of pivotal IBS drug studies – which is also highly encouraging. The only slight divergence appears to be with excluding patients with pain score ≥ 7.5 . We have no concern relative to that, however, as it almost certainly relates to minimizing potential noise given that higher pain scores can often be associated with non-IBS causes.

A "primary endpoint" of IBS-API (i.e. pain measure) is also listed in the clinicaltrials.gov posting – however, that should be considered to only be one of several measures that will be assessed in Study 1 as potential primary endpoints to be incorporated into a pivotal study. Others, as we have indicated in our recent prior updates, will likely include primary endpoints that have been used in recent pivotal studies of IBS drugs including defecation related endpoints (i.e. stool consistency and frequency). We should know more about all of the endpoints under consideration when with future updates to the posting.

The clinical studies will be conducted at Beth Israel Deaconess Medical Center (Harvard teaching hospital) and the University of Michigan. Drs. William Chey and Anthony Lembo are members of BMRA's scientific advisory board and are affiliated with the University of Michigan and Beth Israel Deaconess Medical Center, respectively (and will presumably be the principal investigators at their respective trial sites). As a reminder, Dr. Lembo is Director of the GI Motility Laboratory at the Beth Israel Deaconess Medical Center's Division of Gastroenterology. He is also one of the principal investigators of Biomerica's recently initiated h. pylori clinical study (see below). Dr. Chey Professor of Internal Medicine, Director of the GI Physiology Laboratory, and Director of Medical Services for the Michigan Bowel Control Program at the University of Michigan.

Comment on Primary Endpoint...

We will be very interested to eventually find out what InFoods' pivotal study primary endpoint will be and its similarity (or not) to those used in prescription IBS drugs' pivotal studies (see our Appendix for more detail about primary endpoints).

Given that FDA has already determined InFoods is a non-significant risk diagnostic (i.e. Class I), the efficacy hurdle for U.S. regulatory clearance may be considered to already be set inherently relatively low (which is not uncommon for diagnostics of many conditions and diseases). However, FDA clearance of a diagnostic usually does not portend the commercial potential of that of a drug that also successfully gains U.S. regulatory approval. In other words, while FDA clearance is of obvious importance for InFoods to enter the U.S. market, meaningful physician uptake may require "compelling enough" clinical evidence.

The good news for BMRA is that there are several factors which may mean the "compelling enough" hurdle may not be too difficult to clear – which we discuss below. The choice of primary endpoint in InFoods' clinical studies may have significant influence (perhaps the most influence) on commercial adoption. If InFoods' primary endpoint is the same (or at least similar) to those used in IBS drug pivotal studies, that should allow for an apples-to-apples efficacy comparison between them - and the additional benefit would be that GIs are already versed on them.

But while a similar endpoint might be ideal, as long as whatever measure is chosen has been sufficiently validated (and InFoods demonstrates efficacy), we think there will be demand for InFoods. The fact that InFoods can be used as an addition to IBS drug therapy means that BMRA's diagnostic does not have to demonstrate superiority to current prescription medicine – in this scenario, as long as InFoods can provide incremental benefit to drugs, it should have utility.

“Compelling Enough” May Not Be Tough Given InFoods Appeal to the “3Ps”...

With BMRA's dream-team scientific advisory board presumably guiding InFoods' clinical strategy, we have confidence that whatever endpoint is chosen will have sufficient industry acceptance. Assuming success in the clinical studies, we like the chances for significant adoption of InFoods because of its potential to appeal to the “3Ps” in the healthcare treatment chain – that is, the patient, physician and payer. Here's why;

- IBS Drugs Have Major Drawbacks (see our detailed discussion in Appendix):

- data from pivotal FDA studies of currently available Rx IBS drugs have shown that;
 - they are ineffective for the majority of IBS sufferers
 - they were often barely better than placebo
 - most IBS drugs have a negative effect on IBS that exceeds their benefits¹
- IBS Rx drugs are expensive
 - typical cost-per Rx between \$400 - \$1,000
 - due to the high cost of IBS drugs and the chronic nature of the disease, payers will often require patients to try other, less-expensive, therapies before approving payment
 - lack of efficacy and high cost of drugs drives up total IBS-related healthcare costs as physicians use trial and error approach to try and treat the disease
- IBS drugs can have unpleasant (such as diarrhea) and dangerous (such as risk of pancreatitis) side effects. Some are also not recommended for chronic use
- there is no drug indicated for the treatment of IBS-M

- Pent-up Demand for Better IBS Treatment of Options

- IBS, particularly IBS-D, can be highly detrimental to quality of life
- given that IBS drugs do not provide sufficient relief to most IBS sufferers, physicians and patients are eager for other options
- survey conducted by a 3rd-party research organization found that 70% of physicians surveyed indicated that they would use InFoods

- Reimbursement

- BMRA (as guided by their IBS-expert SAB) believes InFoods is reimbursable under an existing CPT code(s), although they have yet to divulge which one(s)
- we estimate (i.e. guess) that an InFoods diagnosis will charge will be in the range of \$100 - \$300 and a patient can expect to be tested 2 – 3 times per year = \$400 - \$900/year. This is relatively insignificant compared to the ~\$13k/year current average total healthcare cost per IBS-D patient which may further incentivize payers to reimburse for InFoods
- as noted above, payers often require failure of less expensive (i.e. oftentimes non-Rx and not IBS indicated) treatment options prior to authorizing payment for Rx IBS drugs. As such, InFoods may be able to be positioned as a less expensive, first-line option to IBS

Novel H. Pylori Test Clinical Studies Commence Enrollment

The company remains committed to expanding their product menu. While clearly the single major focus is successful development of InFoods, other high-potential projects continue in parallel. Another program, related to a **novel Helicobacter pylori (h. pylori) diagnostic** was the latest to be divulged. BMRA already has several h. pylori tests in their product catalog including ELISA blood antibody tests, a rapid antibody test for the OTC market and a rapid stool-based antigen test for the professional POC market.

H. pylori is a gram-negative bacteria found in the stomach. While it is relatively common – as much as 50% or more of the world's population are infected with the bacteria – most people do not exhibit symptoms. But, of the 15% - 20% that do have a reaction, symptoms can include stomach pain, reflux, nausea and bloating. H. pylori is

¹ Shah, E. and Pimentel, M. (2014), Evaluating the functional net value of pharmacologic agents in treating irritable bowel syndrome. *Aliment Pharmacol Ther*, 39: 973–983. doi:10.1111/apt.12692

associated increased risk of ulcers and is the strongest known risk for developing gastric cancer. Additionally, length of exposure to h. pylori is positively correlated to the risk of gastric cancer. Once diagnosed, standard therapy consists of proton pump inhibitors and certain antibiotics.

In November 2017 BMRA announced that enrollment commenced for the clinical studies for their new and proprietary h. pylori test which “is designed to increase the sensitivity and specificity of H. pylori testing and monitoring of treatment.” The studies are being done in collaboration with the University of Southern California and Vanderbilt University, along with an unnamed European University. Biomerica is ballparking 6 – 12 months for the clinical studies, following which analytical studies will be conducted. BMRA anticipates an eventual application seeking U.S. regulatory clearance will follow FDA’s 510(k) pathway.

Additional details about the study, titled *Specimen Collection Study for H. Pylori Testing in Patients With Dyspepsia*, are listed on clinicaltrials.gov (ID: [NCT02970110](https://clinicaltrials.gov/ct2/show/study/NCT02970110), link: <http://bit.ly/2gRuWLD>). Principal investigators are Dr. Anthony Lembo (Harvard Medical) and Dr. Douglas Morgan (Vanderbilt Medical). Dr. Lembo is also a member of BMRA’s Scientific Advisory Board and a recognized expert in GI disorders. Per clinicaltrials.gov, the study, which will be conducted at a minimum of two sites, will acquire human specimens from patients (n=200) undergoing endoscopy with gastric biopsy for the diagnosis of active h. pylori infection. The biopsy tissue (sampled from the stomach) will be evaluated with histology and rapid urease test (or RUT, a commonly used test which identifies the presence of urease, an enzyme secreted by h. pylori). The study design also notes that a **stool sample will be obtained by the participants prior to undergoing endoscopy** and that “results and specimens will be used in a future clinical trial [i.e. “analytical studies” referenced in BMRA’s PR] of a non-invasive in vitro diagnostic assay for the detection of H. pylori antigen”. **Which we think suggests that BMRA’s novel h. pylori test will be stool-based.**

In April 2018 BMRA contracted with Guardian Angel Research Center to conduct a specimen collection study protocol for h. pylori testing in patients with dyspepsia.

Currently there are several methods to test for the presence of h. pylori. This includes histology, RUT and culture – all of which require invasive biopsy sampling, and non-invasive methods including urea breath test (UBT), serology and stool antigen tests (SAT). There are advantages and disadvantages of each. This includes expense and discomfort of invasive testing and lower accuracy of non-invasive serology testing. Below is a summary (compiled from data of several studies) of the different tests and their relative advantages and disadvantages – the table is from a study by JY Lee and N Kim, published in the January 2015 issue of *Annals of Translational Medicine*.²

Test	Sensitivity	Specificity	Advantages	Disadvantages
Noninvasive				
Serology	76-84	79-90	Widely available, inexpensive	Positive result may reflect previous rather than current infection, not useful after treatment
Urea breath test	>95	>95	High negative and positive predictive values, useful before and after treatment	False-negative results possible in the presence of PPIs or with recent use of antibiotics or bismuth preparations, considerable resources and personnel required to perform test
Stool antigen test	96	97	High negative and positive predictive values, useful before and after treatment	Process of stool collection may be distasteful to patient, false-negative results possible in the presence of PPIs or with recent use of antibiotics or bismuth preparations
Invasive				
Histology	95	99	Excellent sensitivity and specificity, especially with special and immune stains, provides additional information about gastric mucosa	Expensive (endoscopy and histopathology costs), interobserver variability, accuracy affected by PPI and antibiotics use, requires trained personnel
Rapid urease test	90	93	Rapid results, accurate in patients not using PPIs or antibiotics, no added histopathology cost	Requires endoscopy, less accurate after treatment or in patients using PPIs
Culture	58.1	100	Specificity 100%, allows antibiotic sensitivity testing	Variable sensitivity; requires trained staff and properly equipped facilities, expensive

² Ju Yup Lee and Nayoung Kim. Diagnosis of *Helicobacter pylori* by invasive test: histology. *Ann Transl Med*. 2015 Jan; 3(1): 10.

Our Comments: Currently available SAT tests are generally considered highly accurate, although as the table illustrates, there are drawbacks including that the use of PPIs, antibiotics or recent bismuths can affect sensitivity. We will be interested to hear future updates on the progress of the clinical and validation studies and more details about the test under development. Learning more about the novel nature, and how it differs from currently available SATs, will be of particular interest to us. We think that if the test was designed to address one or more of the shortcomings of current SATs – such as, for example, being unaffected by PPIs (with no meaningful compromise to accuracy) or even improving on accuracy, that that may provide for significant differentiation.

Approximately six million non-invasive h. pylori tests are performed each year in the U.S. – if we assume \$100/test, that calculates to a domestic market size of around \$600M. And this is expected to grow given increasing prevalence of h. pylori and a greater shift from direct (i.e. invasive) to non-invasive methods – which could be further catalyzed with the advent of novel technologies (potentially including BMRA's test) addressing some of the drawbacks of currently available tests.

Given BMRA's relatively tiny size (\$34M MC, ~\$6M annual revenue), capturing as little as one-quarter of one percent of the U.S. non-invasive h. pylori market (or ~\$1.5M) would be highly significant for the company. One percent market share could mean doubling of revenue from current levels. We hope to know more about the test in the non-too-distant future which may help in assessing potential competitiveness to currently available diagnostics as well as provide some useful data points for modeling purposes.

VALUATION

We like chances for FDA clearance of InFoods and subsequent commercial demand...

Particularly telling of the desperation and difficulty in developing effective IBS therapies is the low FDA efficacy hurdle used to evaluate IBS drugs and the less than compelling clinical trial data used to (successfully) support FDA approval. Minimal effectiveness and drawbacks of IBS drugs has created what we believe is a relatively low “attractiveness hurdle” and, coupled with a lack of side effects and expected cost-benefit, one that may be easily cleared if InFoods is able to demonstrate only incremental benefit (either alone or to existing therapies).

And in terms of the potential commercial appeal for InFoods, the dearth of sufficiently effective IBS therapies, high related healthcare costs and overall frustration from physicians and patients as a result of lack of better options means high demand already exists for more effective IBS therapies. Couple that with payers’ pushback to reimburse for (relatively expensive) Rx IBS drugs until other (less expensive) options have been tried, and we think appeal to the critical “3Ps” (i.e. patient, physician and payer) in the healthcare treatment chain will be sufficiently satisfied to drive meaningful demand shortly following launch.

InFoods Modeling Assumptions

Market opportunity: IBS is often very difficult to diagnose and symptoms can differ individual-to-individual. This heterogeneity means that the market opportunity for IBS-indicated therapies can also extend to other conditions including symptom-similar diseases (particularly inflammatory bowel diseases) such as Crohn’s, chronic idiopathic constipation (CIC), ulcerative colitis and celiac disease. Our ancillary market size estimates exclude patients comorbid with IBS (i.e. these represent incremental populations). The following is our estimates of the total U.S. market opportunity for InFoods

Total U.S. market opportunity: 58M people

- U.S. IBS total market size: 45M people (18M mild, 16M moderate, 11M severe)
 - IBS-M: 13.5M
 - IBS-C: 13.5M
 - IBS-D: 18M
- U.S. ancillary conditions market size: 13M people
 - Crohn’s: 4M
 - CIC: 2M
 - Ulcerative colitis: 4M
 - Celiac disease: 3M

Initial “reachable” market: while we calculate the total U.S. market for InFoods at approximately 58M people, the initial “reachable” market is likely only a fraction of that as most people with IBS do not visit their doctor for it. According to the International Foundation for Functional Gastrointestinal Disorders, there are only about 3.5M physician visits each year for IBS – which we use as the initial reachable IBS market. For the ancillary markets, we assume the initial reachable market is 30% of the total, or (13M x .30) = 3.8M. However, we also assume that if InFoods can demonstrate incremental efficacy that it will encourage a significantly greater number of people to visit their doctors. We estimate that can increase the size of the reachable market by 15% per year for at least the first five years.

Frequency of InFoods Testing Per Patient: BMRA expects InFoods will be most effective when patients return to be re-tested 2 to 3 times per year which allows for optimizing efficacy through diet re-adjustments. We assume an average of 2.5 visits per patient per year.

Pricing / Margin: we should have a better idea of pricing expectations with more information relative to expected reimbursement. We believe cost (i.e. COGS) of the test and related processing will be relatively inexpensive, particularly with volume, as the technology is largely commoditized and widely available. We currently assume per-test pricing of \$150 and COGS (i.e. material expense and processing) of \$30 and gross profit \$70.

Commercialization Partner: based on our discussions and certain feedback, we think it is likely that BMRA will look to out-license U.S. commercialization, similar to what they did with Telcon in S. Korea. We think out-licensing is also a safer move from a financial standpoint. We hope to know more about potential

commercialization plans over time but, until then, we assume a royalty rate of 15% for the U.S. market (i.e. same rate as Telcon deal).

Market Penetration: given the drawbacks of IBS drugs and pent-up demand for more effective alternatives, we think that if InFoods demonstrates significant incremental clinical benefit at what we expect to be a relatively low cost, it has the potential to see fairly rapid widespread adoption. Additionally, no drugs are indicated to treat IBS-M (~30% of all IBS cases) – which means InFoods uptake among these patients could be particularly robust. We think that InFoods has the potential to capture 10% of the reachable market within the 3rd full-year following launch in the U.S. and 15% by year-5.

Timelines: BMRA is ballparking 9 – 14 months for the (n=180) initial “endpoint-finding” study. While we do not know what to expect in terms of the size or duration of a pivotal InFoods study, our best-guess right now (using pivotal IBS drug trials as a proxy) is somewhere in the range of 300 – 400 patients per arm and 24 months from start to completion (including time for design and any FDA interaction). Assuming another 3 – 6 months for FDA filing and regulatory clearance, results in estimated InFoods U.S. launch around calendar mid-2021 (i.e. early fiscal 2022).

Risk of Unknowns: we are incorporating what could arguably be considered a conservative haircut of 70% to our revenue estimates to account for what remains a fairly sizeable number of significant unknowns. This haircut will be adjusted based on solidifying answers to these unknowns such as, for example, further substantive progress towards validation (from both a regulatory and commercial perspective) of InFoods, reimbursement, market opportunity, intellectual property protection, timelines and U.S. commercialization strategy among others.

United States Only Model					
	Years on market				
	1	2	3	4	5
Assumed physician visits for (in 000s)					
IBS (C,D and M)	3,500	4,025	4,629	5,323	6,122
Ancillary conditions	3,810	4,382	5,039	5,795	6,664
Assumed penetration	0.25%	1.50%	7.00%	12.00%	15.00%
Assumed InFoods tests/patient/yr	2.5	2.5	2.5	2.5	2.5
Assumed revenue/test to distribution ptrnr	\$100.0	\$105.0	\$110.3	\$115.8	\$121.6
Royalty rate	15%	15%	15%	15%	15%
Partner/Distributor revenue (\$000s)	\$ 685.3	\$ 4,965.1	\$ 27,978.3	\$ 57,915.0	\$ 87,415.5
Unknowns haircut	70%	70%	70%	70%	70%
InFoods Revenue to BMRA (\$000s)	\$ 205.59	\$ 1,489.53	\$ 8,393.48	\$ 17,374.51	\$ 26,224.65

Value BMRA at \$3.00/share (base business) + \$4.00/share (InFoods) = \$7.00/share

We use sum of the parts to value BMRA; the base business (everything except InFoods) plus InFoods. We note that we are only modeling assumed U.S. InFoods sales and we do not yet model the novel h. pylori test that recently commenced clinical studies. All of our modeling and valuation-related assumptions will be updated if and when appropriate.

We continue to value the base business using a comparable cohort of five companies of various market capitalizations in the medical diagnostics space with products/services that target the POC and/or clinical lab markets to value BMRA. Based on several metrics, BMRA's base business is valued at approximately \$2.90/share.

Base Business Comparable Multiples

Ticker	P/E (ttm)	P/E (FY1)	P/E (FY4)	P/Book	EV/ EBITDA (ttm)	P/S (ttm)	P/S (FY1)
TRIB	20.9	21.0	13.1	1.2	14.7	N/A	N/A
OXFD	N/A	N/A	N/A	5.2	N/A	3.6	3.0
CEMI	N/A	N/A	0.0	6.6	N/A	4.8	3.8
QDEL	N/A	19.0	35.9	7.7	24.0	6.5	5.3
Average	24.0	32.0	24.7	5.3	24.0	4.9	4.0

FY1, FY4 = forecast year 1, forecast year 4

Value Based On Comp:

	P/E (ttm)	P/E (FY1)	P/E (FY4)	P/Book	P/S (ttm)	P/S (FY1)	AVG
BMRA	N/A	N/A	-\$9.18	\$3.03	\$3.30	\$2.59	\$2.97

Value InFoods at ~\$35M (~\$4.00/share):

We value InFoods separately from that of the base business given the former's significantly greater growth potential. As we outlined above, we think our InFoods assumptions and related outlook/forecast are reasonable. Given the rapid growth rate and steepening revenue inflection that we estimate at approximately years 2023/2024 we think a 11x sales multiple is reasonable, particularly when looking at the ~12x trailing sales multiple Valeant paid for Salix (i.e. Xifaxin) in 2015. Applying 11x to our \$8.4M 2024 forecasted InFoods revenue and discounting back to the present at 15%/year, results in InFoods present value of approximately \$35M, or \$4.00/share. If and when there is attrition of some of the substantive unknowns, our risk discount will similarly reduce and likely result in higher calculated InFoods value.

Our sum of the parts values BMRA at approximately \$7.00/share.

APPENDIX:

InFoods Explained

Important to understand about IBS and why we believe InFoods has so much potential is that while food is implicated as a trigger in exacerbating symptoms, the same foods do not affect every IBS sufferer the same (i.e. different people may have different trigger foods). So it is not as simple as just identifying certain foods and eliminating those from anyone's diet that has the condition. As such, the heterogenous nature of IBS requires a diagnostic (i.e. InFoods) that accounts for varying and different causation between certain foods and an individuals' symptoms, or lack thereof.

Additional details about InFoods have been made public recently – this includes information gleaned from BMRA's patent applications as well as from the company's recent investor presentations.

Below we summarize what we believe are some of the most salient points regarding InFoods;

- blood test to identify certain trigger foods that may cause or exacerbate IBS symptoms
 - will be used only with individuals already identified with IBS symptoms
 - extensive analysis was done to rank the top several dozen foods most associated with exacerbating IBS
 - we expect somewhere in the range of 20 – 25 foods may be included on the initial panel
 - ELISA test quantifies food-specific IgG antibodies (i.e. 'signal scores') from individual IBS patients which are elicited as immune response
 - signal score 'cutpoints' for each food (and gender) were determined by comparing signal scores of IBS patients with those of non-IBS patients. Additional testing and analysis was done to refine these cutpoints, which represent the 90th and 95th percentiles. For each food, IBS subjects with resulting signal scores above these cutpoints are considered 'positive' for that particular food (i.e. that particular food exacerbates IBS)
 - results of the ELISA test provide a simple 'yes' or 'no' result indicating whether a particular person's IBS symptoms are being triggered by each of the foods on the panel
- InFoods is expected to help physicians in guiding treatment protocol including putting the patient on a specific dietary regimen and can be used in combination with IBS drugs. This is different than other IBS tests which only focus on diagnosing presence of the disease. BMRA's test would be the first to both help diagnose IBS and to help guide treatment decisions
- expected to have utility for all forms of IBS (i.e. constipation (C), diarrhea (D) and mixed (M))
- test will be available for use in both the clinical lab and physician office settings. Lab product is the first which they will pursue (regulatory hurdle is likely lower) and POC will follow
- would be reimbursed under existing CPT codes. As reimbursement is critical for maximizing early adoption, availability of payment under existing CPT codes is a significant benefit
- in addition to the significant benefit of already established CPT coding is that it could be expected that patients would be tested more than once (i.e. 2 - 3x) over the course of a year, depending on changes in their diets
- 17 patents are currently pending. In March 2016 BMRA announced International Search Authority reviewed their international method and composition patent claims and found them to be novel and non-obvious (i.e. the claims are valid)
- FDA has indicated that the risk profile of the test would likely not require a Class III (i.e. 'high risk') device designation. This was further supported when in July 2016 BMRA announced that FDA determined InFoods is eligible to pursue 'nonsignificant risk' clinical studies. BMRA expects to apply for the de novo route which allows manufacturers of novel low-risk (Class I and II) products for which there is no predicate to avoid the much costlier and time-consuming PMA route

InFoods[®] Physician's Office (Point of Care) Use (Clinical Lab Version first to be submitted)



InFoods[®] Simple Blood Test

InFoods[®] Test Results

Food	Result
A	+ POSITIVE
B	NEGATIVE
C	NEGATIVE
D	+ POSITIVE
E	NEGATIVE
F	NEGATIVE
G	NEGATIVE
H	+ POSITIVE
I	NEGATIVE
J	NEGATIVE
K	+ POSITIVE
L	NEGATIVE
M	+ POSITIVE
Total # Foods Positive	
5	

Foods are specific Egg + POSITIVE
Blueberry - Negative


◀ 4 ▶

Market considerations related to InFoods: *Living with IBS is hell...*

- “Living with IBS is hell” – plug that into a Google search and it is apparent that the chronic disease symptoms and lack of treatment options leaves IBS sufferers feeling helpless and desperate for more effective options
- Diagnostic cost of IBS in the U.S. is approximately \$10.5B in annual direct costs and over \$30B when including indirect costs
- IBS afflicts as much as 20% of the U.S. population, 25% of Japan, and 22%+ each of China and the U.K
- IBS is a top 10 reason for primary care doctor visits
- IBS is difficult to diagnose and difficult to treat
 - Exact cause of IBS is not known although adverse reaction to certain foods is largely accepted as a significant contributor in many cases
 - Types of foods and food reactions do not appear to be homogenous from patient-to-patient (i.e. cocoa may trigger symptoms in one patient but not another). Therefore it is important to be able to identify which foods trigger symptoms in each individual patient
 - IBS drugs, such as Xifaxan, Linzess, Amitiza, Viberzi and Lotronex (see our discussion below) only treat the symptoms but not the underlying cause, are effective in only about 15% - 20% of IBS sufferers, often provide only temporary and partial relief (in those patients who do show a response) and can have unwelcome side effects
 - While new IBS drugs are in development, including at least one (Trulance) that could gain FDA approval and launch in the coming months, these also only address IBS symptoms and not the underlying cause
 - FDA has a low effectiveness hurdle related to approving IBS drugs – which also speaks to the lack of effective treatments for the disease
 - GI doctors also often prescribe SSRI's off-label, which have shown to help regulate bowel flow – this again, illustrates how limited the treatment options are for IBS
 - Physicians have very limited tools to treat IBS – typical recommendation is for patients to (arbitrarily) begin eliminating certain foods from their diet and/or prescription of symptom-targeting drugs which often fail to provide significant relief, particularly over the long-term
- Win, win, win for patients, physicians and insurers. InFoods could benefit all major stakeholders. Physicians are frustrated with lack of treatment options. Patients feel helpless. Insurers are paying for relatively high cost drugs which do not address the underlying cause and therefore may be chronically prescribed
- Unlike many new medical technologies (drugs, devices and diagnostics) which offer only incremental benefit compared to an existing product and may be geared more towards profit than clinical outcome (and often require a lot of marketing to convince of the ‘benefits’), InFoods could be a pioneer in providing a new level of relief for

the IBS afflicted. And if InFoods can do that, it should require limited initial awareness-building before the test sells itself via demand-pull from physicians and patients

Unmet Need Highlighted By IBS Drugs’ Low Effectiveness, Low FDA Approval Hurdle

The U.S. market for IBS drugs is approximately \$1B+, almost twice the size of what it was only just a few years ago. The market is likely to continue to grow at a rapid pace, largely driven by new drugs that have either recently come to market or that are expected to do so in the near-term. But while the number of drugs available to treat IBS and related expenditures continues to grow, their relative ineffectiveness in providing relief has not significantly changed.

Today, the prescription IBS drug market is mostly concentrated across four products; Linzess (Allergan) and Amitiza (Takeda) for IBS-C, and Xifaxan (Valeant) and Viberzi (Allergan) for IBS-D. No drug is approved in the U.S. for the treatment of IBS-M. All of these gained FDA approval for their respective IBS indications since 2012 and two, Xifaxan and Viberzi, were approved for the U.S. market in May 2015 and launched later that year.

Low FDA Approval Hurdle...

FDA has clearly recognized the seeming futility drugmakers have experienced in their quest to develop a therapy that provides significant and long-term relief of IBS. This is evidenced by the low bar the U.S. regulatory agency has recommended in terms of efficacy endpoints that novel drugs are assessed against in order to determine whether they will be approved for sale. We note that while FDA updated their recommendations for evaluation of novel IBS drugs in 2012 to better assess their effectiveness in addressing symptoms of IBS, the effectiveness hurdle (i.e. improvement vs. placebo) remains low.

The table below describes the pre-updated primary endpoints that have been used in many IBS drug trials. FDA updated their guidelines as they felt that while the prior endpoints could capture the direction of change, they did a poor job of providing useful information of the effect of treatment on sign and symptoms of IBS.

Prior IBS Endpoint Guidelines

Drug and Indication	Primary Endpoint	Questions (Single-Item) Used to Assess Efficacy	Responses
Alosetron — IBS-D ¹	Adequate relief	<i>In the past 7 days, have you had adequate relief of your IBS pain or discomfort?</i>	Binary (Yes/No)
Tegaserod — IBS-C ²	Satisfactory relief	<i>Did you have satisfactory relief of your overall IBS symptoms during the last week?</i>	Binary (Yes/No)
		<i>Did you have satisfactory relief of your abdominal discomfort or pain during the last week?</i>	Binary (Yes/No)
Lubiprostone — IBS-C ³	Subject Global Assessment of Relief (SGA)	<i>Please consider how you felt during the past treatment period in regard to your IBS, in particular your overall well-being, and symptoms of abdominal pain/discomfort and altered bowel habit. Compared to the way you usually felt before entering the trial, how would you rate your relief of symptoms during the past week?</i>	5-Point Likert scale
		<i>How would you rate your relief of IBS symptoms (abdominal discomfort/pain, bowel habits, and other IBS symptoms) over the past week compared with how you felt before you entered the trial?</i>	7-Point Likert scale

SOURCE: US Dept. HHS, FDA, CDER. IBS – Clinical Eval of Drugs. May 2012

The updated recommended guidelines are more defined in terms of assessment of the effect on symptoms and specifically cites a defecation component and an abdominal pain component. For IBS-C drugs, recommendation is that defecation is assessed by stool frequency (number of complete bowel movements per week) while IBS-D drugs

be assessed by stool consistency (based on Bristol Stool Form Scale). The pain component, for both IBS-C and IBS-D, can be based on an 11-point numeric scale.

The table below describes the updated (i.e. 2012) recommended guidelines for primary endpoints of IBS-C and IBS-D investigational drugs. Here we can see that the definition of treatment ‘response’ in these updated guidelines is more directly tied to IBS symptoms (as compared to prior guidelines). But also, the ‘response’ thresholds are arguably meek (and, in the case of the pain measure, potentially still fraught with subjectivity).

Note that ‘response’ related to abdominal pain is a decrease of 30% or more (in weekly average worst pain) in the past 24 hours. Weekly stool consistency ‘response’ is reduction of 50% or more days per week with mushy or watery stools. These endpoints are far from complete reprieve of IBS symptoms and, we think, underscores not only the difficulty in effectively addressing the disease but also, when we look at the substantial revenue generated by and number of prescriptions written for these drugs, how desperate IBS sufferers are for something that will provide even minor and temporary relief.

Current IBS Endpoint Guidelines

Indication	Primary Endpoints	Entry Criteria	Responder Definition
IBS-C	Abdominal Pain Intensity	Abdominal Pain Intensity Weekly average of <i>worst abdominal pain in past 24 hours</i> score of ≥ 3.0 on a 0 to 10 point scale	Abdominal Pain Intensity Decrease in weekly average of <i>worst abdominal pain in the past 24 hours</i> score of at least 30% compared with baseline
	AND Stool Frequency	AND Stool Frequency < 3 CSBMs per week	AND Stool Frequency Increase of 1 or more CSBM per week compared with baseline
IBS-D	Abdominal Pain Intensity	Abdominal Pain Intensity Weekly average of <i>worst abdominal pain in past 24 hours</i> score of ≥ 3.0 on a 0 to 10 point scale	Abdominal Pain Intensity Weekly responder defined as: decrease in weekly average of <i>worst abdominal pain in past 24 hours</i> score of at least 30% compared with baseline
	AND Stool Consistency	AND Stool Consistency At least 2 days per week with at least one stool that has a consistency of Type 6 or Type 7 BSS (see Figure 1 for details)	Daily responder defined as: decrease in <i>worst abdominal pain in the past 24 hours</i> score of at least 30% compared with baseline AND Stool Consistency Weekly responder defined as: decrease at least 50% in the number of days per week with at least one stool that has a consistency of Type 6 or 7 compared with baseline Daily responder defined as: a patient whose stool consistency is less than 5 for all bowel movements on that day or no bowel movement

SOURCE: USDept HHS, FDA, CDER. IBS—Clinical Eval of Drugs. May 2012

Unmet Need For Relief Drives Demand For Drugs, Despite Being Barely Better Than Placebo...

IBS sufferers’ high unmet need for any symptom relief becomes even more obvious when examining primary endpoint outcomes data from the pivotal FDA studies of each of the four prescription drugs that account for 90%+ of the U.S. IBS drug market. In the table below we have summarized efficacy data of the phase III studies used to support FDA approval filings for each of the leading IBS drugs (see Appendix for more details about each compound and the phase III studies). Also included is Trulance, for which an FDA filing for IBS-C is expected to happen in the near term.

Our table also includes estimated current annualized U.S. sales and total prescriptions as well as our comments, all meant to help elucidate the significant demand for IBS drugs despite their relative lack of effectiveness and in many cases, highly unpleasant and even dangerous side effects.

Points of particular interest:

- **Majority of patients did not respond:** response rates ranged from a low of approximately 12% up to 41%

- **Barely more effective than placebo:** response rates as compared to placebo indicate a substantial placebo effect in treatment “response”. While subjectivity of endpoint measures (particularly with ‘legacy’ measures) may play a part (our supposition), the data suggests that none of these drugs are much more effective than placebo. As an example, Linzess, which currently generates ~\$600M in annual U.S. revenue and for which ~2.8M prescriptions are written each year, demonstrated only 7% - 13% superiority on primary efficacy endpoints compared to placebo
- **Drawbacks:** besides cost and relatively low efficacy there are other drawbacks to IBS drugs. This includes side effects which are often unpleasant - such as diarrhea, and which can even be dangerous – Lotronex carries a black box warning related to the risk of potentially serious GI events while Viberzi may cause pancreatitis in people w/o a gallbladder. Some drugs, such as Xifaxan, are not recommended for chronic use while others, such as Amitiza and Lotronex, are only indicated for women due to lack of data for use with men

Indication	Drug	FDA Appr	Treatment Response	Delta vs placebo	Cost*	Annualized U.S.		Comments
						Sales (M)	TRx (M)	
IBS-C	Linzess	2012	12% - 34%	7% - 13%	\$382	\$630**	2.90**	Diarrhea side effect (20%)
	Amitiza	2008	13%	6%	\$380	\$450**	1.50**	Indicated for w omen only as not studied for men
	Trulance	NA	22% - 30%	7% - 12%	NA	NA	NA	Not yet FDA approved
IBS-D	Xifaxan	2015	41%	9%	\$550	\$950***	1.00***	Not for chronic use
	Viberzi	2015	25% - 30%	7% - 14%	\$1,000	\$125	0.15	Abdominal pain (secondary) endpoint not met
	Lotronex	2002	NA	13% - 20%	\$940	\$60	0.04	For w omen only. Black box w arning

*Cost - per Drugs.com. Based on retail price of 30-day supply of lowest dose **Includes CIC and IBS-C

***Includes overt HE and IBS-D

The Low “Attractiveness Hurdle” Should Play In InFoods’ Favor...

While there is not yet enough publicly available information relative to the performance of InFoods to make any informed opinions regarding its clinical effectiveness, the current lack of effective therapies and debilitating symptoms of the disease means that doctors and patients are likely to be receptive to just about anything that has the potential to provide even incremental benefit (whether it be incremental to existing therapies or alone).

We believe that the minimal effectiveness and drawbacks of prescription drugs means that the “attractiveness hurdle” that novel IBS therapies must meet to generate interest from the ~\$1B+ U.S. IBS market is relatively low. That should be particularly true for non-drug, side-effect-free products such as InFoods.

And with InFoods testing expected to be covered with existing CPT codes, coupled with the likelihood of a substantial cost-benefit as compared to IBS drugs, we think the market could be very receptive to the product. We look forward to updates from the company on expected clinical trial design and other details related to validating effectiveness and clinical utility of the test.

IBS Drugs Overview

Xifaxan (Salix/Valeant):

- Xifaxan (rifaximin) is a semisynthetic antibiotic initially approved by FDA (May 2004) for the treatment of traveler’s diarrhea caused by E. coli (200mg), in March 2010 it (550mg) received FDA approval for overt hepatic encephalopathy and in May 2015 it (550mg) received FDA approval for the treatment of IBS-D in adults.
- Recommended dosing is one tablet, 3x/per day for 14 days. Recurrence can be retreated twice at the same dosing regimen
- Method of action is not completely understood, but hypothesized to relate to changes in the bacterial content in the GI tract
- Most commonly reported side effects are nausea (~3%) and increase in ALT (liver enzyme in the blood, 2%)

- Clinical data used to support FDA filing consisted of three double-blind, placebo-controlled phase 3 trials (TARGET 1,2,3).
 - o **Trials 1 and 2:** enrolled 1,258 patients diagnosed (per Rome II criteria) with IBS symptoms of abdominal pain and discomfort. Patients were randomized to Xifaxan (n=624) or placebo (n=634) for 14 days and followed for a subsequent 10-week treatment-free period.
 - o **Primary endpoint** was proportion of patients who achieved adequate relief (based on patients' 'yes' or 'no' response) of IBS symptoms for at least 2 of 4 weeks during the month following 14 days of treatment. Key **secondary endpoint** was proportion of patients who achieved relief of bloating during the same primary evaluation period.
 - o **Results:**
 - **Primary endpoint** of the combined studies: Significantly ($p < 0.001$) more Xifaxan patients (41%) than placebo patients (32%) reported adequate relief of IBS symptoms. Relief from IBS symptoms maintained statistically significant through the entire 3-month study period ($p < 0.001$).
 - **Results on the key secondary endpoint** were similar to that of the primary endpoint with significantly ($p < 0.001$) more Xifaxan patients (40%) than placebo patients (30%) reporting adequate relief of IBS-related bloating.
 - o **Trial 3** was powered to assess effectiveness of Xifaxan in cases of recurrence. 636 patients who did not meet a composite endpoint in TARGET 1,2 of relief of abdominal pain and loose stools were randomized in TARGET 3 for retreatment and received Xifaxan (n=328) or placebo (n=308) for another 14 days, followed by a 4 week treatment-free period. **Results** showed that significantly ($p = 0.02$) more Xifaxan (33%) patients achieved symptom relief as compared to placebo patients (25%)

Primary endpoint	Proportion of Patients with Response				Difference	P-value
	Xifaxan	%	Placebo	%		
<i>Weekly global IBS symptoms</i>						
Target 1	126/309	40.8%	98/314	31.2%	9.6%	0.01
Target 2	128/315	40.6%	103/320	32.2%	8.4%	0.03
Combined	254/624	40.7%	201/634	31.7%	9.0%	< 0.001
Key secondary endpoint						
<i>Weekly IBS-related bloating</i>						
Target 1	122/309	39.5%	90/314	28.7%	10.8%	0.005
Target 2	129/315	41.0%	102/320	31.9%	9.1%	0.02
Combined	251/624	40.2%	192/634	30.3%	9.9%	< 0.001

Viberzi (Allergan):

- Viberzi (eluxadoline) received FDA approval May 2015 for the treatment of IBS-D in adults. Launched in U.S. December 2015
- Eluxadoline targets opioid receptors (it is a μ - and κ -opioid receptor agonist and δ -opioid receptor antagonist) in the portion of the nervous system that govern the GI tract, thereby reducing bowel contractions and related pain.
- Recommended dosing is one tablet, 2x/day and can be taken as long as doctor recommends
- Most commonly reported side effects are constipation, nausea and abdominal pain
- In March 2017 FDA issued a warning against its use with that patients w/o gallbladder as it could cause serious pancreatitis
- Clinical data used to support FDA filing consisted of two Phase 3 double-blind, placebo controlled studies
 - o Combined studies enrolled 2,425 patients (study 1 = 1,280, study 2 = 1,145) diagnosed with IBS-D (Rome III criteria) randomized to either 75mg or 100mg Viberzi (n=1,616) twice daily or placebo (n=809)
 - o Patients treated over 26 weeks
 - o Primary endpoint was simultaneous improvement in worst abdominal pain score by $\geq 30\%$ as compared to baseline and improvement in stool consistency (based on Bristol Stool Scale) on at least 50% of the days within a 12-week time interval (for FDA) and through 26 weeks (for EMA)

- Secondary endpoints included improvement in the defined pain score at 12 weeks and improvement in the defined stool consistency score at 12 weeks
- **Results:**
 - 12 weeks on composite (primary FDA) endpoint (pain and stool):
 - Study 1: 24% (75mg) to 25% (100mg) of Viberzi patients vs. 17% of placebo patients met composite endpoint, indicating statistically significant improvement favoring treatment arm of 7% - 8% ($p < 0.05$)
 - Study 2: 29% to 30% of Viberzi patients vs. 16% of placebo patients met composite endpoint, indicating statistically significant improvement favoring treatment arm of 13% - 14% ($p < 0.001$)
 - 26 weeks on composite (primary EMA) endpoint (pain and stool):
 - Study 1: 23% to 29% of Viberzi patients vs. 19% of placebo patients met composite endpoint. The 10% difference to the 100mg arm was statistically significant ($p \leq 0.014$) while the 4% difference to the 75mg arm was not
 - Study 2: 30% to 33% of Viberzi patients vs. 20% of placebo patients met composite endpoint. The 13% - 14% difference was statistically significant ($p \leq 0.014$)
 - 12 weeks on abdominal pain (secondary) endpoint:
 - Study 1: 42% to 43% of Viberzi patients vs. 40% of placebo patients reported reduction in abdominal pain. The 2% - 3% difference was not statistically significant
 - Study 2: 48% to 51% of Viberzi patients vs. 45% of placebo patients reported reduction in abdominal pain. The 3% - 6% difference was not statistically significant
 - 12 weeks on stool consistency (secondary) endpoint:
 - Study 1: 30% to 34% of Viberzi patients vs. 22% of placebo patients reported improvement in stool consistency. The 8% - 12% difference was statistically significant
 - Study 2: 36% to 37% of Viberzi patients vs. 21% of placebo patients reported improvement in stool consistency. The 15% - 16% difference was statistically significant
 - **Rescue loperamide: % of patients requiring rescue loperamide during 26-week treatment phase**
 - Study 1: 25% of Viberzi vs. 28% of placebo
 - Study 2: 29% of Viberzi vs. 35% of placebo

	Study 1			Study 2		
	VIBERZI 100mg twice daily n=426	VIBERZI 75mg twice daily n=427	PBO n=427	VIBERZI 100mg twice daily n=382	VIBERZI 75mg twice daily n=381	PBO n=382
Composite¹ Response over 12 weeks						
Responder rates	25%	24%	17%	30%	29%	16%
Treatment difference	8% ²	7% ⁴		13% ³	13% ³	
95% CI (%)	(2.6, 13.5)	(1.4, 12.2)		(7.5, 19.2)	(6.8, 18.5)	
Composite Response over 26 weeks						
Responder rates	29%	23%	19%	33%	30%	20%
Treatment difference	10%	4%		13%	10%	
95% CI (%)	(4.7, 16.1)	(-1.0, 9.9)		(6.4, 18.8)	(4.2, 16.4)	
Abdominal Pain Response Improved $\geq 30\%$ over 12 weeks						
Responder rates	43%	42%	40%	51%	48%	45%
Treatment difference	4%	3%		6%	3%	
95% CI (%)	(-3.0, 10.2)	(-3.8, 9.4)		(-1.3, 12.8)	(-4.3, 9.8)	
BSS < 5 Response over 12 weeks						
Responder rates	34%	30%	22%	36%	37%	21%
Treatment difference	12%	8%		15%	16%	
95% CI (%)	(6.3, 18.2)	(2.1, 13.8)		(8.4, 21.0)	(9.7, 22.4)	

¹ Composite= Simultaneous improvement of Worst Abdominal Pain (WAP) by $\geq 30\%$ and Bristol Stool Score (BSS)

< 5 on the same day for $\geq 50\%$ of days over the interval

² P=0.01

³ P<0.001

⁴ P=0.05

SOURCE: Viberzi label, Actavis. Accessdata.fda.gov

Linzess (Allergan):

- Linzess (linaclotide) is a peptide agonist of guanylate cyclase 2C (GC-C) approved by FDA in 2012 for the treatment of adults with IBS-C as well as for the treatment of chronic idiopathic constipation (CIC)
- Recommended dosing is one tablet daily
- Most commonly reported side effects are diarrhea (20%), abdominal pain (7%) and flatulence
- Method of action: linaclotide stimulates secretion of intestinal fluid resulting in acceleration of GI transit and reduction on abdominal pain
- Clinical data used to support FDA approval for IVS-C consisted of two double-blind, randomized, placebo-controlled phase 3 studies
 - o Combined studies enrolled 1,604 patients (study 1 = 800, study 2 = 804) diagnosed with IBS-C (Rome II criteria) randomized to either Linzess (n=806) or placebo (n=798)
 - o Patients treated for 12 weeks
 - o Primary endpoint: there were four primary endpoints:
 - In at least 9 of the first 12 weeks of treatment patient had to have;
 - Combined responder endpoint (9 of 12 weeks): $\geq 30\%$ reduction from baseline in abdominal pain, ≥ 3 complete spontaneous bowel movements (CSBM) and an increase of ≥ 1 CSBM from baseline, all in the same week
 - $\geq 30\%$ reduction from baseline in abdominal pain
 - ≥ 3 complete spontaneous bowel movements (CSBM) and an increase of ≥ 1 CSBM from baseline in the same week
 - Combined responder endpoint (6 of 12 weeks): In at least 6 of the first 12 weeks of treatment patient had to have $\geq 30\%$ reduction from baseline in abdominal pain and an increase of ≥ 1 CSBM from baseline, all in the same week
 - o **Results:**
 - Combined responder 9 of 12 weeks:
 - Study 1: 12% of Linzess vs. 5% of placebo met endpoint

- Study 2: 13% of Linzess vs. 3% of placebo met endpoint
- Abdominal pain 9 of 12 weeks:
 - Study 1: 34% of Linzess vs. 27% of placebo met endpoint
 - Study 2: 39% of Linzess vs. 20% of placebo met endpoint
- CSBM 9 of 12 weeks:
 - Study 1: 20% of Linzess vs. 6% of placebo met endpoint
 - Study 2: 18% of Linzess vs. 5% of placebo met endpoint
- Combined responder 6 of 12 weeks:
 - Study 1: 34% of Linzess vs. 21% of placebo met endpoint
 - Study 2: 34% of Linzess vs. 14% of placebo met endpoint
- Differences between treatment and placebo arms on all endpoints were statistically significant

At least 9 of first 12 weeks

	Trial 1			Trial 2		
	LINZESS 290 mcg (N=405)	Placebo (N=395)	Treatment Difference [95% CI]	LINZESS 290 mcg (N=401)	Placebo (N=403)	Treatment Difference [95% CI]
Combined Responder* (Abdominal Pain and CSBM Responder)	12.1%	5.1%	7.0% [3.2%, 10.9%]	12.7%	3.0%	9.7% [6.1%, 13.4%]
Abdominal Pain Responder* (≥ 30% Abdominal Pain Reduction)	34.3%	27.1%	7.2% [0.9%, 13.6%]	38.9%	19.6%	19.3% [13.2%, 25.4%]
CSBM Responder* (≥ 3 CSBMs and Increase ≥1 CSBM from Baseline)	19.5%	6.3%	13.2% [8.6%, 17.7%]	18.0%	5.0%	13.0% [8.7%, 17.3%]

At least 6 of first 12 weeks

	Trial 1			Trial 2		
	LINZESS 290 mcg (N=405)	Placebo (N=395)	Treatment Difference [95% CI]	LINZESS 290 mcg (N=401)	Placebo (N=403)	Treatment Difference [95% CI]
Combined Responder* (Abdominal Pain and CSBM Responder)	33.6%	21.0%	12.6% [6.5%, 18.7%]	33.7%	13.9%	19.8% [14.0%, 25.5%]

* Primary Endpoint, ** Secondary Endpoints
 Note: Analyses based on first 12 weeks of treatment for both Trials 1 and 2
 CI = Confidence Interval

SOURCE: Linzess label. Allergan

Amitiza (Takeda):

- Amitiza (lubiprostone) received initial FDA approval in 2006 for the treatment of chronic idiopathic constipation in adults and in 2008 received FDA approval for the treatment of women with IBS-C
- Amitiza is a bicyclic fatty acid that acts on GI epithelial cells to promote secretions which helps to soften the stool and increase motility
- Recommended dosing for IBS-C is orally, twice daily
- Most common side effects are nausea (8%), diarrhea (7%) and abdominal pain (5%)
- Clinical data used to support FDA approval for IBS-C consisted of two double-blind, placebo-controlled phase 3 studies (Note: FDA guidelines for evaluating novel IBS drugs changed in 2012; i.e. after Amitiza rec'd FDA approval)
 - o Combined studies enrolled 1,154 patients (92% of which were female) with IBS-C and received Amitiza 2x/day or placebo for 12 weeks
 - o Primary endpoint compared the proportion of "Overall responders" in each arm based on 7-point global relief questionnaire. "Monthly responder" was defined as reporting "significantly relieved" for at least 2 weeks of the month or at least "moderately relieved" in all four weeks of the month. "Overall responders" were those that were "monthly responders" for at least 2 of the 3 months of the study.
 - o Results:
 - Study 1: 14% of Amitiza patients vs. 8% of placebo patients were "overall responders"
 - Study 2: 12% of Amitiza patients vs. 6% of placebo patients were "overall responders"
 - In both studies the difference between treatment and placebo was statistically significant

Lotronex (Prometheus):

- Lotronex (Alosetron) originally received FDA approval in February 2002 for the treatment of women with IBS-D but the manufacturer voluntarily removed it from the market later that year due to reports that use of it had been associated with life-threatening adverse effects including serious intestinal damage and severely obstructed bowels. In 2002 FDA approved an sNDA, allowing for restricted marketing of Lotronex only for women with severe diarrhea-predominant IBS. Less than 5% of IBS is considered severe. The revised labeling also includes a black box warning related to potentially serious GI adverse events
- Lotronex acts as an agonist to receptors of the GI-related nervous system, helping to increase water absorption and slow GI motility
- Recommended dosing is orally, twice per day for 12 weeks
- Most commonly reported side effects are constipation (29%), abdominal pain (7%) and nausea (6%)
- Clinical data: Lotronex has been studied in three clinical studies in women with severe diarrhea-predominant IBS, including two studies (i.e. studies 1,2) with patients with bowel urgency $\geq 50\%$ of days and one study (i.e. study 3) with patients with 1 or more of the following: frequent and severe abdominal pain or discomfort, frequent bowel urgency or fecal incontinence, disability or restriction of daily activities due to IBS
 - o **Results;**
 - Studies 1,2: Lotronex patients had 13% to 16% (statistically significant) greater increase compared to placebo patients in the median percentage of days with bowel urgency control. In addition, 50% of Lotronex patients had bowel urgency no more than 1 day in the last week compared to 29% of placebo patients. Also, 12% of Lotronex patients had bowel urgency no more than 2 days per week in any of the 12 treatment weeks compared to 1% of placebo patients
 - Study 3: Lotronex dosed at three different doses; 0.5mg/day, 1mg/day or 1mg 2x/day. All Lotronex groups had significantly greater number of responders (43% to 51%) versus placebo (31%) based on 7point GIS scale at 12 weeks

IN LATE-STAGE DEVELOPMENT

Trulance (Synergy):

- Trulance (plecanatide) received FDA approval for the treatment of adults with chronic idiopathic constipation in January 2017. It has also completed two phase 3 trials for the treatment of IBS-C, an indication for which Synergy filed for FDA approval (sNDA) of in March 2017. FDA approval and launch could happen in 1H 2018
- Similar to Linzess, Trulance is a GC-C agonist. It is a 16 amino acid peptide which acts to stimulate increased intestinal fluid and accelerate GI transit.
- Recommended dosing for IBS-C is orally once/day
- Clinical studies: Data from two (3mg and 6mg doses) phase 3 studies in IBS-C was announced in May 2017.
 - o Combined studies enrolled 2,189 patients (study 1 = 1,135, study 2 = 1,054) with IBS-C who were randomized to Trulance (n = 1,456) or placebo (n = 733)
 - o Patients treated once daily for 12 weeks
 - o Primary endpoint is percentage of patients who are "overall responders", defined as those patients that simultaneously experience $\geq 30\%$ reduction in worst abdominal pain and increase of ≥ 1 CSBM from baseline, in the same week, for at least 50% of the 12 treatment weeks
 - o **Top-line results**
 - Study 1 (n = 1,135): 21.5% of Trulance 3mg and 24.0% of Trulance 6mg patients vs. 14.2% of placebo patients met primary endpoint. The difference between placebo and both Trulance dose cohorts was statistically significant (p=0.009 for 3mg, p<0.001 for 6mg)
 - Study 2 (n = 1,054): 30.2% of Trulance 3mg and 29.5% of Trulance 6mg patients vs. 17.8% of placebo patients met primary endpoint. The difference between placebo and both Trulance dose cohorts was statistically significant (p<0.001) for 3mg, p<0.001 for 6mg)
 - Adverse events: the most common adverse events in the IBS-C phase 3 studies were diarrhea (~4%)

Scientific Advisory Board

Dr. Douglas Drossman

Dr. Drossman is a veteran of over 50 FDA clinical trials and is the current president of the Rome Foundation, a leading international organization that provides support and guidance in the diagnosis and treatment of functional gastrointestinal disorders. The Rome Foundation is responsible for creating the Rome process, and the criteria developed through this process are the most widely employed in clinical trials related to IBS.

Dr. Lin Chang

Director, Digestive Health and Nutrition Clinic UCLA GI Fellowship Training Program and Professor, Digestive Diseases/Gastroenterology. Dr. Chang has significant experience with pharmaceutical and healthcare companies as she has been an advisor or consultant to over 32 major pharmaceutical companies, including GlaxoSmithKline, Novartis, Merck, Allergan, Takeda, Salix, Synergy, Johnson & Johnson, Entera Health, and Ardelyx. Her experience with the FDA includes serving on the Gastrointestinal Drugs Advisory Committee of the FDA 2005-2010 (Chair 2009-2010) and again 2015-2019 as well as working as a FDA Special Government Employee from 2009-2013.

Dr. William Chey

Professor of Internal Medicine, Director of the GI Physiology Laboratory, and Director of Medical Services for the Michigan Bowel Control Program at the University of Michigan. Dr. Chey has also worked with many major pharmaceutical and healthcare companies as an advisor or consultant, including Entera Health, Ironwood, Nestle, Proctor and Gamble, Salix, and Takeda. He is the Chair of the U.S. Scientific Advisory Board at SmartPill Corporation and a member of the Clinical Advisory Board at Synthetic Biologics.

Dr. William Whitehead

Director of the Center for Functional GI & Motility Disorders and Professor of Medicine Adjunct Professor of OB-GYN at University of North Carolina School of Medicine. He has worked on 29 NIH grants (19 as principal investigator, 10 as co-investigator) and has been continuously funded by NIH since 1977. On behalf of the International Foundation for Functional Gastrointestinal Disorders, he organized two international consensus conferences on the treatment of fecal incontinence (1999 and 2002) and led a workshop on design of treatment trials for pharmaceutical companies, academic investigators, the NIDDK, and the FDA for 8 years. As a consultant, he has worked with large pharmaceutical companies on clinical trial design, including Takeda, Sucampo, Ironwood, Forest, Ono, and McNeil.

Dr. Anthony Lembo

Director of the GI Motility Laboratory at the Beth Israel Deaconess Medical Center's (BIDMC) Division of Gastroenterology in Boston, MA and as an Associate Professor of Medicine at Harvard Medical School. Dr. Lembo completed his residency and GI Fellowship at UCLA Medical Center. He is an accomplished expert in afflictions of the gastrointestinal tract and IBS. He divides his time between clinical medicine and research at Beth Israel Deaconess Medical Center in Boston. He has authored numerous original clinical studies and other research articles related to IBS.

Strategic Advisory Board

Ned Barnholt

Currently chairman of the KLA-Tencor Corporation and serves on the board of directors of eBay and Adobe. He is the former chairman, president, and chief executive officer of Agilent Technologies, a leading company in life sciences, diagnostics and applied chemical markets. Mr. Barnholt led the Agilent Technologies spin-off of Hewlett-Packard Company which broke records as the largest initial public offering (IPO) in Silicon Valley history at the time of the IPO (US \$2.1 billion).

Harry Leider, MD

Until recently, Dr. Leider served as Chief Medical Officer (CMO) and Group Vice President of Walgreens. As the CMO at Walgreens, he provided executive leadership for health analytics, clinical program development, and clinical sales solutions. Dr. Leider also served as the senior clinical spokesperson for the company and routinely represented Walgreens with leaders in government, managed care, health systems, and other healthcare organizations. As Chief Medical Officer, he led a cross-functional department of over 40 professionals that conducted over 60 health outcomes studies that demonstrated the value of Walgreens programs and services. He also had leadership responsibility for a team that evaluated over 200 emerging healthcare technologies to provide information about potential M & A and partnership opportunities. Notably, Dr. Leider also directed the clinical design for a digital health programs that rewards 3 million Walgreens customers for taking steps to improve their health.

Prior to Walgreens, Dr. Leider was the Chief Medical Officer of Ameritox which was the nation's largest specialty lab serving clinicians who treat chronic pain and behavioral health conditions. He was responsible for the development of all provider support and research programs, and played a key role supporting sales efforts directed at providers and payors. Prior to his role at Ameritox, Dr. Leider held C-suite positions in several population health and payor companies.

FINANCIAL MODEL

Biomerica Inc.

	2018 A	Q1E	Q2E	Q3E	Q4E	2019 E	2020 E	2021 E	2022 E
InFoods TOTAL	\$0.0	\$0.0	\$0.0	\$0.0	\$0.0	\$0.0	\$0.0	\$0.0	\$205.6
<i>YOY Growth</i>	-	-	-	-	-	-	-	-	-
Legacy TOTAL	\$5,564.0	\$1,327.9	\$1,437.8	\$1,675.2	\$1,526.4	\$5,967.2	\$6,386.6	\$6,882.8	\$7,089.3
<i>YOY Growth</i>	-3.9%	-8.0%	-10.9%	21.7%	35.1%	7.2%	7.0%	7.8%	3.0%
Total Revenues	\$5,564.2	\$1,327.9	\$1,437.8	\$1,675.2	\$1,526.4	\$5,967.2	\$6,386.6	\$6,882.8	\$7,294.9
<i>YOY Growth</i>	-3.9%	-8.1%	-10.9%	21.8%	35.0%	7.2%	7.0%	7.8%	6.0%
Cost of Goods Sold	\$3,809.8	\$896.3	\$938.9	\$1,088.9	\$995.2	\$3,919.3	\$4,017.2	\$4,239.8	\$4,376.9
Gross Income	\$1,754.4	\$431.6	\$498.9	\$586.3	\$531.2	\$2,048.0	\$2,369.4	\$2,643.0	\$2,918.0
<i>Gross Margin</i>	31.5%	32.5%	34.7%	35.0%	34.8%	34.3%	37.1%	38.4%	40.0%
SG&A	\$1,837.8	\$431.6	\$473.0	\$539.4	\$488.4	\$1,932.4	\$2,031.0	\$2,168.1	\$2,268.7
<i>% SG&A</i>	33.0%	32.5%	32.9%	32.2%	32.0%	32.0%	31.8%	31.5%	31.1%
R&D	\$1,398.4	\$518.2	\$531.6	\$544.0	\$558.4	\$2,152.2	\$3,820.0	\$2,944.0	\$2,550.0
<i>% R&D</i>	25.1%	39.0%	37.0%	32.5%	36.6%	36.1%	59.8%	42.8%	35.0%
Operating Income	(\$1,481.8)	(\$518.2)	(\$505.7)	(\$497.1)	(\$515.7)	(\$2,036.7)	(\$3,481.5)	(\$2,469.1)	(\$1,900.8)
<i>Operating Margin</i>	-26.6%	-39.0%	-35.2%	-29.7%	-33.8%	-34.1%	-54.5%	-35.9%	-26.1%
Total Other Income (Expense)	\$47.7	\$6.2	\$2.1	\$10.9	\$8.1	\$27.3	\$30.0	\$25.0	\$40.0
Pre-Tax Income	(\$1,434.0)	(\$512.0)	(\$503.6)	(\$486.2)	(\$507.6)	(\$2,009.4)	(\$3,451.5)	(\$2,444.1)	(\$1,860.8)
Tax expense (benefit)	\$31.8	\$0.0	\$0.0	\$0.0	\$0.0	\$0.0	\$0.0	\$0.0	\$0.0
<i>Tax Rate</i>	-2.2%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%
Net Income	(\$1,465.8)	(\$512.0)	(\$503.6)	(\$486.2)	(\$507.6)	(\$2,009.4)	(\$3,451.5)	(\$2,444.1)	(\$1,860.8)
<i>YOY Growth</i>	61.3%	113.5%	72.9%	24.5%	-10.9%	37.1%	71.8%	-29.2%	-23.9%
<i>Net Margin</i>	-26.3%	-38.6%	-35.0%	-29.0%	-33.3%	-33.7%	-54.0%	-35.5%	-25.5%
EPS	(\$0.17)	(\$0.06)	(\$0.06)	(\$0.05)	(\$0.06)	(\$0.22)	(\$0.37)	(\$0.26)	(\$0.19)
<i>YOY Growth</i>	56.8%	104.2%	65.0%	21.0%	-12.2%	30.4%	67.0%	-30.8%	-25.6%
Diluted Shares O/S	8,570	8,955	8,987	9,035	9,071	9,012	9,270	9,480	9,700

Brian Marckx, CFA

HISTORICAL ZACKS RECOMMENDATIONS



DISCLOSURES

The following disclosures relate to relationships between Zacks Small-Cap Research (“Zacks SCR”), a division of Zacks Investment Research (“ZIR”), and the issuers covered by the Zacks SCR Analysts in the Small-Cap Universe.

ANALYST DISCLOSURES

I, Brian Marckx, CFA, hereby certify that the view expressed in this research report accurately reflect my personal views about the subject securities and issuers. I also certify that no part of my compensation was, is, or will be, directly or indirectly, related to the recommendations or views expressed in this research report. I believe the information used for the creation of this report has been obtained from sources I considered to be reliable, but I can neither guarantee nor represent the completeness or accuracy of the information herewith. Such information and the opinions expressed are subject to change without notice.

INVESTMENT BANKING AND FEES FOR SERVICES

Zacks SCR does not provide investment banking services nor has it received compensation for investment banking services from the issuers of the securities covered in this report or article.

Zacks SCR has received compensation from the issuer directly or from an investor relations consulting firm engaged by the issuer for providing non-investment banking services to this issuer and expects to receive additional compensation for such non-investment banking services provided to this issuer. The non-investment banking services provided to the issuer includes the preparation of this report, investor relations services, investment software, financial database analysis, organization of non-deal road shows, and attendance fees for conferences sponsored or co-sponsored by Zacks SCR. The fees for these services vary on a per-client basis and are subject to the number and types of services contracted. Fees typically range between ten thousand and fifty thousand dollars per annum. Details of fees paid by this issuer are available upon request.

POLICY DISCLOSURES

This report provides an objective valuation of the issuer today and expected valuations of the issuer at various future dates based on applying standard investment valuation methodologies to the revenue and EPS forecasts made by the SCR Analyst of the issuer’s business. SCR Analysts are restricted from holding or trading securities in the issuers that they cover. ZIR and Zacks SCR do not make a market in any security followed by SCR nor do they act as dealers in these securities. Each Zacks SCR Analyst has full discretion over the valuation of the issuer included in this report based on his or her own due diligence. SCR Analysts are paid based on the number of companies they cover. SCR Analyst compensation is not, was not, nor will be, directly or indirectly, related to the specific valuations or views expressed in any report or article.

ADDITIONAL INFORMATION

Additional information is available upon request. Zacks SCR reports and articles are based on data obtained from sources that it believes to be reliable, but are not guaranteed to be accurate nor do they purport to be complete. Because of individual financial or investment objectives and/or financial circumstances, this report or article should not be construed as advice designed to meet the particular investment needs of any investor. Investing involves risk. Any opinions expressed by Zacks SCR Analysts are subject to change without notice. Reports or articles or tweets are not to be construed as an offer or solicitation of an offer to buy or sell the securities herein mentioned.