

## Mast Therapeutics (MSTX - \$ 0.42)

### 2Q15: One Step Closer to EPIC Study Top-line Results – Anticipated in 1Q16

Yesterday, MSTX reported 2Q15 financial results with a net loss of (\$10.2MM) vs. Laidlaw (\$9.0MM) and the Street (\$9.1MM) estimates. Net loss per share equated to (\$0.06), which is identical to that of Laidlaw and the Street. The company ended 2Q15 with cash of ~\$43MM, enough to support its operation deep into 2016, in our opinion.

- Patient enrollment of EPIC study is on-track.** MSTX reported that patient recruitment of the EPIC trial is on track with 70% of patients already enrolled (vs. >50% in April 2015) and top-line results expected in 1Q16. The average age of patients is 14/15 years old. Vepoloxamer remains clinically the most advanced and broadly applicable SCD therapy in development to date.
- Multiple elective supportive studies could strengthen the label.** MSTX will conduct a supportive study evaluating vepoloxamer in renal impaired adult patients to enrich the safety database for assessing body's capacity of removing excess drug if over-dosed. This study along with QTC, repeating dosing (EPIC-E) and an upcoming tissue oxygenation study are all elective (not required by the FDA), and they could potentially strengthen the label of vepoloxamer if the drug is approved.
- Pipeline developments.** MSTX plans to start a vepoloxamer in chronic heart failure Phase II trial in Sep. 2015 with top-line results potentially available in 2016. This would be a randomized, double-blind, two-arm, placebo-controlled and 150-patient study. The objectives of the trial are to examine safety and efficacy, which include the drug's effect on biological markers of cardiac injury (troponin), wall stress (NT-proBNP), and other clinical outcomes. MSTX will terminate the vepoloxamer in acute limb ischemia (ALI) Phase II trial and start a stroke Phase II study in 2016 (possibly 2H16). The sole purpose of the ALI trial is to provide some data so it could advance to the stroke study. We also expect the preliminary data of an investigator-sponsored trial that evaluates AIR001 in WHO Group 2 pulmonary hypertension (PH) patients associated with left heart disease potentially be available in 2H15. Two AIR001 studies are ongoing.
- Action.** We are reiterating our Buy rating and our \$2.50 target price based on peer comparable probability adjusted DCF analyses to reflect the continued execution of corporate developments.

#### Earnings Estimates: (per share)

(Dec)	1Q	2Q	3Q	4Q	FY	P/E
<b>FY-15E</b>	-0.06A	-0.06A	-0.06	-0.06	-0.25	N.A.
<b>FY-14A</b>	-0.06	-0.06	-0.06	-0.05	-0.23	N.A.
<b>FY-13A</b>	-0.12	-0.09	-0.05	-0.06	-0.28	N.A.
<b>FY-12A</b>	-0.09	NA	-0.07	-0.08	-0.33	N.A.

Source: Laidlaw & Company estimates

Healthcare/Biotechnology

Ticker:	<b>MSTX</b>
Rating:	<b>Buy</b>
Price Target:	<b>\$ 2.50</b>

#### Trading Data:

Last Price (08/13/2015)	\$ 0.42
52-Week High (9/8/2014)	\$ 0.69
52-Week Low (8/10/2015)	\$ 0.38
Market Cap. (MM)	\$ 65
Shares Out. (MM)	164

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- **Balance sheet strengthened.** MSTX announced it has entered into a loan and security agreement with Hercules Technology to borrow up to \$15MM in two tranches. MSTX excised the first tranche of \$5MM upon entry into the agreement on August 11, 2015. The second tranche of \$10MM is available through the end of 2015, provided that vepoloxamer and AIR001 programs achieve certain clinical development milestones.

**Table 1: Estimated and reported 2Q15 results**

<b>2Q15 Estimates and Reported Results</b>			
<b>(\$,000)</b>	<b>Laidlaw Estimate</b>	<b>Actual</b>	<b>Consensus</b>
<b>Total revenue</b>	<b>\$0.0</b>	<b>\$0.0</b>	<b>\$0.0</b>
<b>Total op. profit (loss)</b>	<b>(\$9,055)</b>	<b>(\$10,181)</b>	<b>(\$9,130)</b>
R&D	\$6,163	\$7,734	
SG&A	\$2,862	\$2,410	
<b>EPS</b>	<b>(\$0.06)</b>	<b>(\$0.06)</b>	<b>(\$0.06)</b>
Net income (loss)	(\$9,018)	(\$10,151)	(\$9,107)

Source: Bloomberg, SEC filings and Laidlaw and Co.

### Anticipated milestones in 2015 and beyond

Product	Indication	Event	Timing	Importance
MST-188	Vaso-occlusive crisis (VOC) in sickle cell disease (SCD)	Periodical updates on Phase III trial progress	2015	***
		Completion of Phase III (EPIC) study	4Q15	***
		Report of Phase III study top-line results	1Q16	****
		Potential NDA filing	2H16	***
		Potential approval	2H17	****
	Stroke	Potentially start Phase II trial	2016	***
	Chronic heart failure	Potentially start Phase II trial	3Q15	***
Potentially report Phase II top-line results		4Q16	****	
AIR001	PH associated with heart failure with preserved ejection fraction (HFpEF),	Potentially start investigator-sponsored Phase II study	2015	***
		Report preliminary results	2H15	****
R&D day			Oct. 7, 2015	***

\*\*\*\* / \*\*\*\*\* Major catalyst event that could impact share price very significantly while \*\*\* event is more informative

Source: Laidlaw & Company and company presentation

## Major Risks

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**Risks of clinical study failure could have a major impact on MSTX share value.** Despite an encouraging prior Phase III study outcome and potentially favorable trial design of the ongoing EPIC Phase III study, risks still exist that MST-188 might not receive approval by the FDA if the pivotal study does not meet its endpoints. Given that the majority of upside for MSTX shares is currently based on the assumption that a positive EPIC study outcome could lead to MST-188 approval before its commercial potential can be realized; a failure of the EPIC study would have a significant negative impact on MSTX share value.

**Commercial success of the MST-188 in resolving vaso-occlusive crisis in sickle cell disease (SCD) patients is less predictable.** Although we recognize that the substantial unmet medical need of shortening the VOC in SCD patients, we cannot fully predict the market acceptance and potential revenue ramp up for the product; as the actual clinical performance will play an important role even if the product is approved. In addition, we cannot fully foresee the market dynamic if competitors also enter the market since such dynamic could be affected by multiple factors. Together, commercial performance of MST-188 may not meet expectations, and if so, MSTX share value could also be impacted negatively.

**Lack of patent protection could make MST-188 vulnerable if competitors develop method to generate a me-too product that might not require complete clinical studies but as generic.** Although the production processes of MST-188 are protected on several fronts, including proprietary technology, trade secrets and know-how, it remains possible that other competitors could develop similar or alternative processes to produce a similar, or even better, product like MST-188. As such, the company might not enjoy the competitive edge and potentially damage MST-188's commercial outlook.

**Limited product diversity could increase overall risk.** Given the nascent stage of the corporate development, majority of the product pipeline value mainly resides on MST-188 in SCD development. The second potential pipeline product, AIR001 in pulmonary arterial hypertension and additional indications (acute limb ischemia, stroke and acute heart failure), potentially could be addressed by MST-188 remains in very early development stage. As such, we believe the company at the current stage has very limited diversification potential in its product pipeline.

**Additional financing could dilute shareholder value.** Although the company ended 3Q14 with ~\$43MM cash, MSTX could potentially need more financial resources going forward if they want to expand and further develop its pipeline and/or commercialize MST-188 in SCD. Should the product not receive FDA approval or product revenue does not reach expectations, the company might need to increase its financial resources, which includes issuing new equity to raise additional cash. Under such a scenario, the share value of existing shareholders could be diluted.

Figure 1: Income Statement

Mast Therapeutics – Income Statement														
(\$'000)	2012	2013	2014	1Q15	2Q15	3Q15E	4Q15E	2015E	2016E	2017E	2018E	2019E	2020E	2021E
<b>Revenue</b>														
MST-188 revenue	0	0	0					0	0	28,253	86,796	167,886	252,822	331,386
Net sales	0	0	0	-	-	-	-	0	0	0	0	0	0	0
Licensing revenue	0	0	0	-	-	-	-	0	0	0	0	0	0	0
Grant revenue	0	0	0	-	-	-	-	0	0	0	0	0	0	0
Total revenue	0	0	0	-	-	-	-	0	0	28,253	86,796	167,886	252,822	331,386
Costs of goods	0									2,543	7,812	15,110	22,754	29,825
Research and development	8,088	12,902	19,436	6,042	7,734	7,889	8,046	29,711	31,791	27,022	25,671	26,441	27,235	28,052
Selling, general and administrative	7,519	8,518	9,487	3,578	2,410	2,453	2,490	10,932	11,260	12,273	13,377	14,581	15,748	16,850
Marketing and sales										20,000	23,000	24,380	25,599	26,367
Transaction-related expenses	(70)	80	271	-	-	0	-	0	0	0	0	0	0	0
Depreciation and amortization	90	40	84	30	37	37	37	141	141	141	141	141	141	141
<b>Total Operating Expenses</b>	15,628	21,539	29,279	9,650	10,181	10,379	10,574	40,784	43,191	61,979	70,001	80,653	91,476	101,235
Operating Incomes (losses)	(15,628)	(21,539)	(29,279)	(9,650)	(10,181)	(10,379)	(10,574)	(40,784)	(43,191)	(33,726)	16,794	87,232	161,346	230,151
Reduction of fair value of warrants	0	0	0	-	-	0	-	0	0	0	0	0	0	0
Investment income	74	60	67	17	30	17	17	81	89	98	108	119	130	143
Interest expense	0	0	0	0	0	0	-	0	0	0	0	0	0	0
Other income/(expense), net	(5)	(1)	511	17	0	20	23	60	2	(20)	24	(27)	(27)	(27)
Loss before cumulative effect of change in accounting principle	(15,559)	(21,480)	(28,701)	(9,616)	(10,151)	(10,342)	(10,534)	(40,643)	(43,100)	(33,648)	16,926	87,324	161,449	230,268
Cumulative effect of change in accounting principle	0	0	0	-	0	0	-	0	0	0	0	0	0	0
<b>Income before tax</b>	(15,559)	(21,480)	(28,701)	(9,616)	(10,151)	(10,342)	(10,534)	(40,643)	(43,100)	(33,648)	16,926	87,324	161,449	230,268
Tax	0	0	0	-	-	0	-	0	0	0	(6,263)	(32,310)	(59,736)	(85,199)
<b>Net Income (Loss)</b>	(15,559)	(21,480)	(28,701)	(9,616)	(10,151)	(10,342)	(10,534)	(40,643)	(43,100)	(33,648)	10,664	55,014	101,713	145,069
Net Income (Loss) Applicable to Common Shareholders	(15,559)	(21,480)	(28,701)	(9,616)	(10,151)	(10,342)	(10,534)	(40,643)	(43,100)	(33,648)	10,664	55,014	101,713	145,069
Net Earnings (Losses) Per Share—Basic and Diluted	(\$0.33)	(\$0.28)	(\$0.23)	(\$0.06)	(\$0.06)	(\$0.06)	(\$0.06)	(\$0.25)	(\$0.25)	(\$0.20)	\$0.06	\$0.32	\$0.59	\$0.83
Shares outstanding—basic	47,641	76,586	122,409	159,459	162,128	163,128	164,128	162,211	169,211	170,211	171,211	172,211	173,211	174,211
Shares outstanding—diluted	47,641	76,586	122,409	159,459	162,128	163,128	164,128	162,211	169,211	170,211	171,211	172,211	173,211	174,211
<b>Margin Analysis (% of Sales/Revenue)</b>														
Costs of goods									9%	9%	9%	9%	9%	9%
R&D	NA	NA	NA	NA	NA	NA	NA	NA	NA	96%	30%	16%	11%	8%
MG&A	NA	NA	NA	NA	NA	NA	NA	NA	NA	114%	42%	23%	16%	13%
Operating Income (loss)	NA	NA	NA	NA	NA	NA	NA	NA	NA	-119%	19%	52%	64%	69%
Pretax	NA	NA	NA	NA	NA	NA	NA	NA	NA	-119%	20%	52%	64%	69%
Tax Rate										37%	37%	37%	37%	37%
Net Income	NA	NA	NA	NA	NA	NA	NA	NA	NA	-119%	12%	33%	40%	44%
<b>Financial Indicator Growth Analysis (YoY%)</b>														
Licensing revenue								0%	0%	0%	0%	0%	0%	0%
Grant revenue								0%	0%	0%	0%	0%	0%	0%
Total Revenue	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	207%	93%	51%	31%
R&D	40%	60%	51%	41%	60%	46%	63%	53%	7%	-15%	-5%	3%	3%	3%
SG&A	5%	13%	11%	58%	2%	0%	4%	15%	3%	9%	9%	8%	7%	7%
Marketing and sales											15%	6%	5%	3%
Operating Income (Losses)	17%	38%	36%	41%	41%	32%	44%	39%	6%	-22%	-150%	419%	85%	43%
Net Income	17%	38%	34%	51%	42%	31%	44%	42%	6%	-22%	-132%	416%	85%	43%
EPS	-31%	-14%	-16%	-1%	1%	-1%	27%	7%	2%	-22%	-132%	413%	84%	42%
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Source: Bloomberg LP; Company reports; Laidlaw & Company estimates

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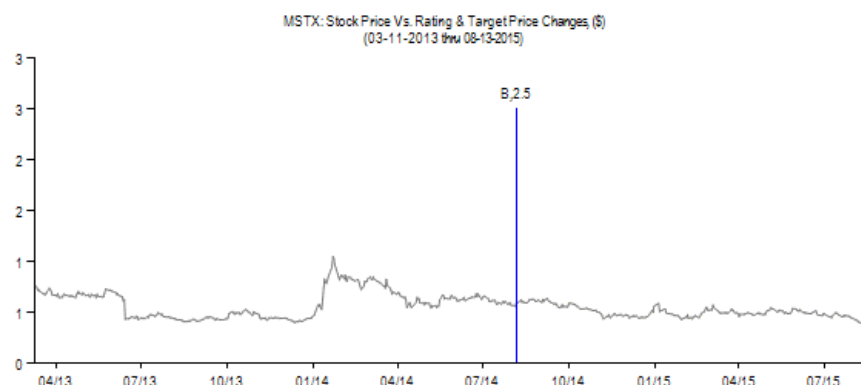
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#### Rating and Price Target Change History



#### 3 Year Rating Change History

Date	Rating	Closing Price (\$)
08/06/2014	Buy (B)	0.60

#### 3 Year Price Change History

Date	Target Price (\$)	Closing Price, (\$)
08/06/2014	2.50	0.60

Source: Laidlaw & Company

Created by: Blue-Compass.net

Laidlaw & Company Rating System*		% of Companies Under Coverage With This Rating	% of Companies for which Laidlaw & Company has performed services for in the last 12 months	
			Investment Banking	Brokerage
<b>Strong Buy (SB)</b>	Expected to significantly outperform the sector over 12 months.	0.00%	0.00%	0.00%
<b>Buy (B)</b>	Expected to outperform the sector average over 12 months.	75.86%	31.03%	6.90%
<b>Hold (H)</b>	Expected returns to be in line with the sector average over 12 months.	3.45%	0.00%	0.00%
<b>Sell (S)</b>	Returns expected to significantly underperform the sector average over 12 months.	0.00%	0.00%	0.00%

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