QUERY RECEIVED FROM KAVAN DIGISH PANDIT [digish.pandit@gmail.com] / APURVA JAYANTKUMAR MEHTA [APURVAJMEHTA@GMAIL.COM]

No. of Shares held: 700 shares / 1,020 shares

Date: 16.04.2025

Sl No.	Query	Management response			
	SMS LIFESCIENCES				
1)	Break up of Sales (9 Months Ended 31 st December 2024) –	= Domestic - 67.64%			
	Domestic and Exports – Percentage	\equiv Exports - 33.36%			
2)	Geography Wise Break Up of Exports – Percentage				
		% to Exports			
		Other Countries 17% RUSSIA 3% GERMANY 3% HONG KONG 4% SPAIN 7% TURKEY 7% BRAZIL 8% 9%			
3)	What is the Company's plan to increase Exports to USA, Europe	Company is actively forging strategic partnerships, expanding its product portfolio			
	and other global markets?	& on boarding new customers as part of our focused efforts to drive export growth.			
4)	What is your Growth Strategy for the Domestic Market?	Company has established long-term relationships with its domestic customers and is actively working to enhance business opportunities in APIs, intermediates, and			
		contract manufacturing. Additionally, it has a long-term strategy in place to drive			
		sustainable revenue growth and capacity utilization.			

5)	Which are the New Products introduced by the Company.	Majorly Trazadone, Furosemide, Cimetidine, Minoxidil etc			
6)	 Please engage in the following activities: Investor Presentation Press Release on Quarterly Results Calls and Meetings with Analysts and Investors Detailed Annual Report for FY25 with a lot of information on the Company 	Company is currently evaluating the feasibility of releasing an "Investor Presentation" and/or "Press Release" on an <u>annual basis</u> , as part of its efforts to enhance transparency and engage effectively with stakeholders. Company remains fully compliant with all statutory disclosure requirements, including circulation of the Annual Report within the prescribed timelines laid down			
	by SEBI (Listing Obligations and Disclosure Requirements) Regulations, 2015. MAHI DRUGS (material subsidiary)				
7)	What is the Company's plan for Exports to USA, Europe and other global markets ?	2-3 products are awaiting approval now in this year and commercialization of at1-3 APIS/Intermediates is targeted every year			
8)	What is the progress on your Strategic Tie Up with ChemWerth (USA) ?	Strategic investment from ChemWerth Inc was completed with equity infusion of ₹45 cr against 40% stake in Mahi Drugs in the year 2021-22 & 2022-23. Several products marketed by ChemWerth are currently at the development and validation stage. Commercial supplies of these products are expected to commence from the next financial year onwards.			

CAUTIONARY STATEMENT:

This response may be "forward-looking statements" within applicable Laws, Rules and Regulations. Actual results could differ from those expressed or implied.

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Query received from Dhruv Bajaj <u>dhruvbajaj184@gmail.com</u> No. of Shares held: 80 shares Date: 06.03.2025

Sl No.	Query	Management response
	TOPIC – RANITIDINE	
1)	As per my reading, Ranitidine was removed from the essential drugs list, so is the	Removal from essential drugs list has nothing to do with ban. It
	drug banned in countries like the USA & Europe?	is related to list of few drugs for price and supply monitoring by
2)	What is our sales mix between Ranitidine & non-Ranitidine-based products in our	Government of India. Currently there is no sale in US but there
	revenues, & the geographical mix of Ranitidine?	are sales in India, Latin America, Asia, CIS, China.
3)	 What's your take on the recent controversy surrounding Ranitidine & are we gearing ourselves for any potential ban in the Indian market? Are our manufacturing facilities fungible to shift from Ranitidine to a lesser impurities-based drug like Famotidine which we are already producing? Or are they not substitutable in nature & any potential ban will hit our topline? 	Current studies have not conclusively proven that Ranitidine directly causes cancer in humans. NDMA, a known carcinogen, consumed at levels exceeding the acceptable daily limit (0.096 mcg), is associated with cancer risks. It is prevalent in common foods and water. It is safe drug in the market for many decades.
4)		
5)	We have repeatedly mentioned in the past that Ranitidine has lower margins hence any adverse impact in that biz will hit our topline more the bottom line, however, as per my discussion with (employee name) from your team, we have done some strong backward integration in Ranitidine leading to higher margins. Therefore, do we believe that this 15 % OPM% is sustainable or will they revert to a mean of 8-12%?	Our volumes on Ranitidine have been stable for last few years. Our Ranitidine sales are currently 45%. Our sales other than Ranitidine are increasing YoY and will ensure any adverse impact is taken care off, though we don't
6)	Players like Strides Pharma exited the market because of the withdrawal of Ranitidine in 2020, however, we have scaled up our business post-2022, so what has led to this stark performance & why have other players not adopted the same strategy Vs completely withdrawing the industry? Has the consolidation in the industry led to higher margins?	foresee any issue in the future. Further, we have no comments on the strategy of other companies. For the FY 2022 there were other campaigns of other customers along with ranitidine, which contributed to the top line. For other Financial years, Quantities of Ranitidine are intact except between October 2019 to September 2020.

7)	As per the credit rating reports, the share of ranitidine in our revenues has increased	However, the increase and decrease of revenues are due to	
	substantially in the past 2 years post the resolution of the NDMA controversy,	variance in price realisation of Ranitidine in line with raw	
	however, if I look at the consolidated numbers from Fy19 till FY24, our bottom line	material prices. We are anticipating the sustainability of	
	has remained flat throughout the period\So can you please share what allowed us to	operating margin at 12 – 15%.	
	maintain the topline during the FY19-22 period of the NDMA issue & why have our		
	topline not expanded substantially post FY23 since the resolution of the issue?		
8)	Is it a fair assessment that our standalone facilities are running at near full capacity	Capacity utilisation is up to 70% and there is scope for another	
	utilisations & incremental sales growth from FY26 will come from the subsidiary? Or	15 to 20% depending upon product mapping and there is	
	do we have plans to debottleneck/incur new capex in the standalone business?	revenue and profitability increase expected from SMS	
		Lifesciences India Limited as well. Also Subsidiary will	
		contribute to increase of revenues/profitability in the next 2-4	
		years as well.	
	Mahi Drugs (Material Subsidiary)		
9)	In my previous conversation with (employee name), he mentioned the reason behind	It is conveyed that the facility is utilised for validation batches	
	the losses in the subsidiary was the maintenance shutdown in Q2 & one can expect a	and their clearances but not for maintenance. Q3 Mahi Drugs	
	ramp-up of operations in Q3, however, the same hasn't materialized. So what might	has reported reasonable sales and have eliminations which did	
	be the reasons behind this?	not reflected in consolidation.	
10)	Congratulations, on receiving the USFDA approvals for the subsidiary. Can you please	Thank you so much for your kind words on the successful	
	give some information regarding the drugs that we will be targeting from this entity	USFDA. We have 2-3 products awaiting approval in this year	
	& the target market for our APIs/intermediates?	and commercialization of 3-4 APIS/Intermediates is targeted	
		every year.	
11)	By when can we expect the proper ramp-up of Mahi drugs given the recent	We are anticipating promising years from FY 2026-27.	
	regulatory approvals? Do we already have some API fillings in that subsidiary which		
	we can immediately commercialise in FY26?		

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	allocation policy going forward?	years.
13)	Our last credit rating report came in FY24, what is the reason behind the lack of any	Credit rating availed and it is available in public domain.
	new credit ratings for the company? & can we expect an updated rating in the coming	
	quarters?	
14)	Given the receipt of USFDA approval & the recent name change, do we plan to release	We are considering the feasibility of investor presentation at
	investor presentations or press releases explaining our quarterly results, if not	appropriate time.
	quarterly conference calls?	
15)	What is the contribution of the top 3 drugs in our total revenues & what is the	We are targeting 10-15% growth for the next few years.
	guidance for coming years directionally given the new molecules potentially to be	
	launched in the subsidiary?	
16)	What will be the incremental capex requirements in the subsidiary in the coming 2-3	We will add two new API lines (one in this year and one in next
	years, or is the existing block sufficient to meet our growth aspirations for next 2-3	year). We will be exporting from Mahi Drugs apart from
	years? & would we continue to do the backward integration work for our partner in	backward integration for the parent company. In coming years
	the subsidiary or will it focus entirely on exporting API's?	the consolidated revenues will increase.

Cautionary Statement:

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