

Investor Presentation

2023 1Q Report

Prepared in accordance with China Accounting Standards



Performance Highlights and Financial Review

Strengths and Key Growth Drivers

Pharmaceutical

Med Tech

Healthcare Services

Appendix

Performance Highlights and Financial Review

Performance Highlights (1/2)

Launched Product



Serplulimab Injection (PD-1)

- 2023Q1 revenue RMB250 million
- Approved for MSI-H, sqNSCLC, ES-SCLC in **Chinese Mainland**
- SCLC was granted with Orphan-drug **Designation** from FDA and EC
- The MAA of ES-SCLC was accepted by the EMA

Trastuzumab Injection (HER2)



- 2023Q1 domestic sales RMB540 million (+66.7% YoY)
- The BLA for breast cancer and metastatic gastric cancer indications was accepted by the FDA

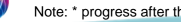


Keverprazan Hydrochloride

Launched in Chinese Mainland for the treatment of duodenal ulcer (DU) and reflux esophagitis (RE)

Product Pipeline

- **FS-1502 (HER2-ADC)** for HER2-positive locally advanced or metastatic breast cancer initiated Ph3 clinical trial in Chinese Mainland
- FCN-159 (MEK small molecule) * for histiocytic tumors, HLX208 (BRAF V600E Inhibitor)* for adult Langerhans Cell Histiocytosis (LCH) and Erdheim-Chester Disease (ECD) with BRAF V600E mutation have been granted with the Breakthrough Therapy **Designation** by the NMPA
- Completed the enrollment of the Ph3 clinical trial of 13-valent pneumococcal conjugate vaccines (multivalent combinations) * in Chinese Mainland
- The **BLA of RT002 (DaxibotulinumtoxinA)** *for the ٠ temporary improvement of moderate to severe glabellar lines in adults caused by corrugator supercilii and/or procerus muscle activity was accepted by the NMPA



Performance Highlights (2/2)

Internationalization

- Collaborated with Syneos Health and initiated Serplulimab injection (PD-1) prelaunch in the U.S.
- Controlled subsidiary Sisram established new direct sales team in Dubai and proposed to control the brand and channels of "PhotonMed" through M&A to achieve direct sales of energy-based devices in China in 1Q2023; established new direct sales team in the UK in 2022; the direct sales revenue accounts for 66% of the 2022 revenue
- Controlled subsidiary Gland Pharma fully acquired Cenexi and entered into Europe-based CDMO by the end of April 2023
- Building localized manufacturing capacities in Africa Côte d'Ivoire

Collaboration

 License-out: granted the exclusive development and commercialization rights for rituximab injection in 16 emerging markets in Asia and Africa to Boston Oncology, to further enhance the accessibility

Performance Highlights



1Q23 Financial Review

Key Financials (RMB million)	1Q23	1Q22*	YoY	Expense Structure	1Q23	1Q22*	Key Indicators	1Q23	2022
Revenue ¹	10,871	10,385	4.7%	Gross Margin	51.3%	43.6%	Cash and bank balances (RMB million)	15,473	16,241
Net profit attributable to shareholders	987	457	116.2%	Selling and Distribution	23.1%	19.9%	Net asset attributable to shareholders	45,393	44,582
Net profit after one-off gain/loss ¹	919	801	14.8%	Administrative	9.1%	7.6%	(RMB million)	1.08	1.06
Net operating cash flow	873	856	2.1%	R&D	8.9%	7.8%		1.00	1.00
R&D Expense	969	810	19.7%	Finance	2.4%	1.0%	Quick ratio	0.85	0.85
Basic EPS (RMB/share)	0.37	0.18	105.6%	Gross Margin minus Selling and Distribution	28.2%	23.6%	Debt-to-asset ratio	48.7%	49.5%

Note 1: the increase of revenue and net profit after one-off loss was caused by the combined impact of: 1) improved product portfolio and increased sales from new launches in the past few years including Serplulimab Injection (PD-1), Trastuzumab Injection(HER2), Avatrombopag Maleate and Azvudine; 2) decreased overseas sales of third party personal protective products for COVID-19; 3) decreased sales of Comirnaty (mRNA COVID-19 vaccine)

Note 2: nonrecurring gain RMB68 million (+RMB412 million YoY), mainly due to market fluctuations of BNTX and other stocks held by the Group results approximately RMB344 million one-off loss in 2022Q1(Fosun Pharma sold all BNTX stake by the end of 2022)

Note 3: the increase of gross margin minus selling and distribution was mainly due to: 1) improved product portfolio and increased sales from new launches in the past few years; 2) the decreased overseas sales of third party personal protective products for COVID-19 with lower gross margins

Note 4: Due to the business consolidations for enterprises under common control in 2022, the Group made retrospective adjustments to the comparative financial information in accordance with the PRC Accounting Standards for Business Enterprises.

2022 Financial Review

Key Financials (RMB million)	2022	2021	YoY	Expense Structure	2022	2021	Key Indicators	2022	2021
Revenue	43,952	39,011	12.7%	Gross Margin	47.3%	48.1%	Cash and bank balances (RMB million)	16,241	10,317
Net profit attributable to shareholders	3,731	4,729	-21.1%	Colling and Distribution	20.0%	22.29/			
Net profit after one-off gain/loss	3,873	3,277	18.2%	Selling and Distribution	20.9%	23.3%	Net asset attributable to shareholders (RMB million)	44,582	39,196
Net operating cash flow	4,218	3,938	7.1%	Administrative	8.7%	8.3%		4.00	
R&D Expenditure	5,885	4,978	18.2%	R&D	9.8%	9.8%	Current ratio	1.06	1.04
R&D Expense	4,302	3,837	12.1%				Quick ratio	0.85	0.85
Basic EPS (RMB/share)	1.43	1.85	-22.7%	Finance	1.5%	1.2%			
Dividend Payout Ratio (Subject to approval by the shareholders)	30%	30%	-	Gross Margin minus Selling and Distribution	26.4%	24.8%	Debt-to-asset ratio	49.5%	48.2%

Note :

Note: nonrecurring loss RMB142 million (-RMB1,593 million YoY), mainly due to market fluctuations of BNTX and other stocks held by the Group; the net effect of BNTX disposal and fair value changes results approximately RMB1 billion one-off loss; realized RMB3,731 million (-21.10% YoY) net profit attributable to shareholders in 2022

<u>The decrease of Gross Margin was mainly due to:</u> 1) the lower gross margins on overseas sales of third party personal protective products for COVID-19; 2) the unit price increase of some core products due to the increase in labor costs and raw materials; 3) but the GM of Pharma business increased by 2.96 pct due to the continuous optimized product structure

 The decrease of selling and distribution rate was caused by the combined impact of: 1) continuously strengthen the control of sales expanse; 2) the decreased selling and distribution rate of volume based purchasing products;3) spend on market development and sales team for new launches in the past few years including Serplulimab injection (PD-1) Note : the increase of cash and bank balances was mainly due to the raised RMB4.48 billion from nonpublic placement of A-Shares in July 2022. The raised fund is for 1) innovative drug clinical trials, license-in and launch; 2) construction manufacturing base for API and formulation; 3) replenishment working capital



Strengths and Key Growth Drivers

Strengths

Constructing internationally competitive asset structure and building organizational capabilities with forward-looking industry insights and operation experiences



Strengths of the Group

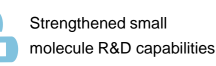
Industry background



Upgraded Innovative Pipeline & System Development - R&D Strategy

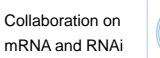
Core Technology Platform

Small Molecule, Antibody/ADC, RNA, Cell Therapy



Established R&D capabilities of novel antibody including monoclonal antibody, bispecific antibody and ADC

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03

Strengthening CAR-T leadership and expanding to immune cell therapy

Core Therapeutic Areas

3 strategic care therapeutic areas and other areas of interest



Oncology Immunity Chronic Disease (liver disease, metabolism, kidney disease)



Other areas of interest: rare disease, antiinfection, cardiovascular, etc.



Building a dynamic and efficient global R&D system which is resultoriented and innovation-driven

Core R&D System and Capabilities

- Efficient and comprehensive "end-to-end" R&D capabilities from project management to market launch
- Clinical value-oriented drug innovation, FIC+BIC accounts for over 50% of the pipeline products
- Accelerated the R&D of competitive product with dynamic evaluation



Upgraded Innovative Pipeline & System Development - Core Products

Launched Core Product

Core Product Pipeline

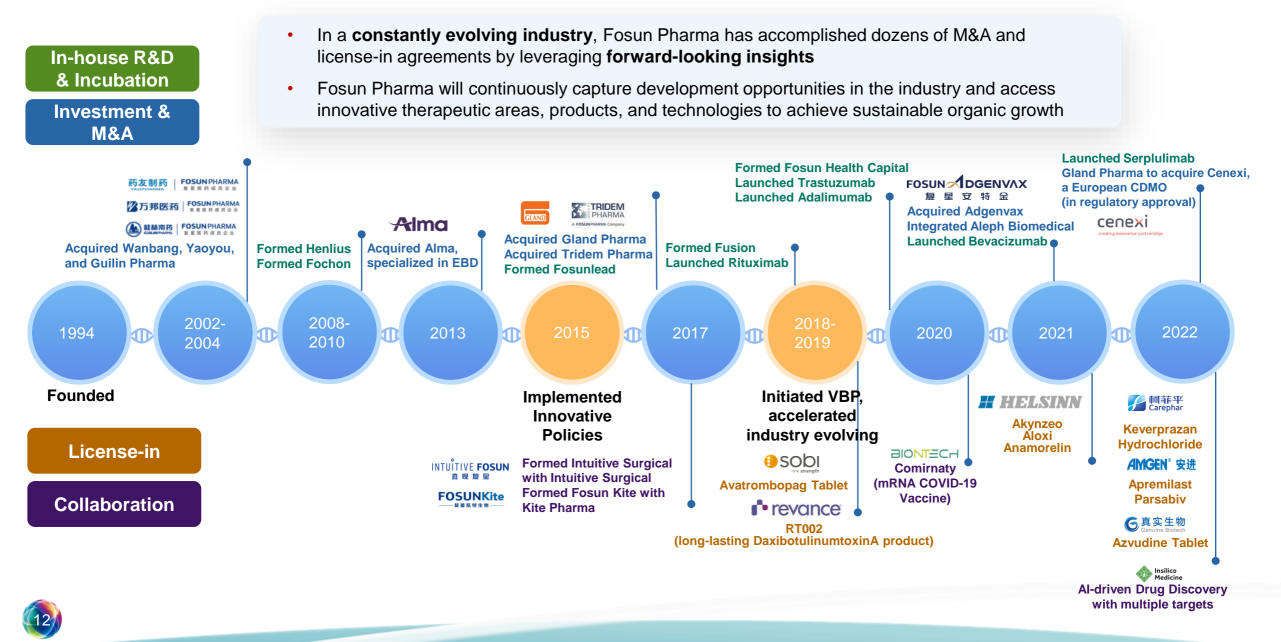
	Serplulimab (PD-1) MSI-H, sqNSCLC, ES-SCLC	Ejilunsai (CAR-T) Third-line LBCL	NDA				Ejilunsai (CAR-T) Second-line LBCL ng-lasting botulinum toxin) GL, CD	
	Rituximab (CD20) Lymphoma, RA	Trastuzumab (HER2) Breast Cancer		Avatrombopag Maleate ITP		Etelcalcetide HPT		
	Netupitant and Palonosetron		Ph3	Serplulimab injection (Neo-/adjuvant treatment of ga	astric cancer	Ē	nor (NHE3 small molecule) ESRD-HD, IBS-C	
Innovative Products	Chemo-induced nausea and vomiting	Azvudine COVID-19 Treatment		FCN-437 (CDK4/6) Breast Cancer	Breast Ca			
	Avatrombopag Maleate	Apremilast	Ph2	Hematological malignancies	FCN-338 (Bcl-2) Hematological malignancies; R/R BCL FCN-159 (MEK small molecule)		208 (BRAF V600E) lid tumor, LCH, ECD ET-26	
	CLDT	Psoriasis	Other	Type I Neurofibron	na	Anesthesia FKC-889 (CAR-T)		
	Antimalarial Series Including Artesunate	Keverprazan Hydrochloride – Chinese Mainland	Pivotal Studies	Keverprazan Hydrochloride DU, RE	e - Giodai	F	MCL	
	Anti-malarial	Duodenal Ulcer, Reflux Esophagitis						
Vaccines	mRNA COVID-19 Vaccine Hong Kong, Macau, Taiwan regions COVID-19 Prevention	Bivalent mRNA COVID-19 Vaccine Hong Kong, Macau, Taiwan regions COVID-19 Prevention		Ph3 13-Valent Pneumocou Conjugate Vaccine Pneumococcal Disease Pr	e	Ph1 C	Valent Pneumococcal Conjugate Vaccine Soccal Disease Prevention	
Vaccilles	Human Rabies Vaccine (Vero Cells) <i>Rabies Prevention</i>	Influenza Vaccine		Ph3 Freeze-dried Human Rabies Vaccine (Vero Cells) Rabies Prevention		Ph3 4-Valent Influenza Vaccine Influenza Prevention		
Generics	27 generic drugs / indic Chinese Mainland / Hong K		Filed 30 generic o R&D pipeline by the er					
s N	ote: last update on 28 th April 2023							



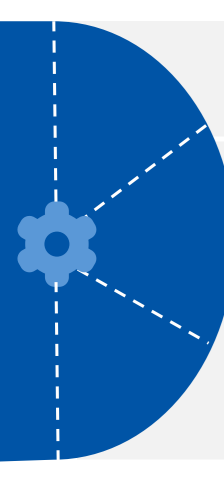
Note:

Non-oncology drugs

Access to Opportunities Through In-house R&D, Incubation, Strategic M&A and Collaboration



Lean Management System



Integrating API and formulation manufacturing and focusing on key pipelines

- Building a regionalized manufacturing center around Xuzhou Area, vertically integrating Sino API facility with Xuzhou formulation facility to achieve intensive production capacity, covering multiple dosages and disease areas
- Chongqing facility and Changde facility have completed the first stage construction; Sino API facility and Xuzhou formulation facility have completed the tech transfer and validation for the first batch. The increased capacity will support future commercial manufacturing

Fosun Ecosystem/Entrepreneurship System, lean management and improvement of daily management system

- Achieved closed-loop procurement management through SRM system, promoting standardization, digitalization and intelligence business
- Improved R&D and clinical trials management, cost control and R&D team synergy by implementing an end-to-end R&D management platform based on in-house developed INNOX digital platform
- Incremental FES projects in 2022 covering quality, cost, efficiency, cycle time, R&D, etc.

Commercialization integration and optimization to control sales expenses and improve sales efficiency

- Commercialization team matches with current product portfolio; 6,000 people in pharmaceutical commercialization team covers oncology and non-oncology areas, OBM broad market team, OTC, online channels and teams in Africa, India and the U.S.
- Strengthening effective control of sales expenses, with the growth rate of sales expenses lower than the growth rate of revenue; the sales expense ratio was 20.87% in 2022 (-2.46 pp YoY)
- Key products cost reduction and efficiency improvement, preparing for procurement and transforming marketing model



Global Operation (1/2)

 Gland Pharma to fully acquire Cenexi for up to EUR210 million and to enter into Europe-based CDMO with localized manufacturing capability

Europe

Africa

- Established 5 regional distribution hubs; the Kenya distribution hub has passed the on-site inspection of the ICRC
- Constructing the Côte d'Ivoire Industrial
 Park with R&D, manufacturing and
 distribution capabilities, localizing products
 manufacturing and distributing in the future

O India

Gland Pharma **Dexrazoxane** for Injection is approved in Chinese Mainland in February 2023; filed several other products in Chinese Mainland

China

- Focusing on **complex** injectables and expanding to biologics CDMO
- Fully acquired Cenexi and entered into Europe-based CDMO by the end of April 2023

The U.S. Generic Drugs: collaborated with 5 major wholesalers and 16 GPOs. Rapid growth in sales of formulations

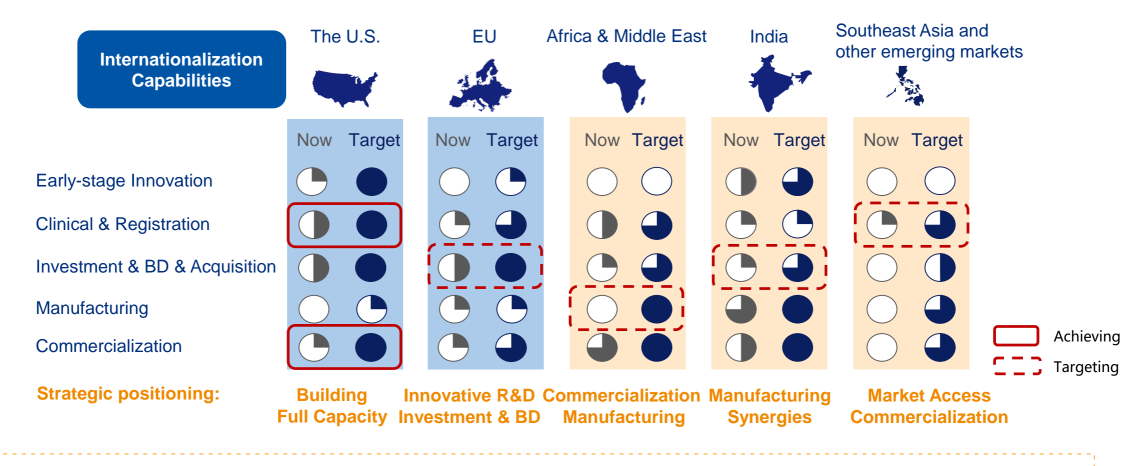
Innovative Drugs:

- 11 combination therapies with Serplulimab Injection (PD-1) are in global multi-center clinical trials; initiated head-to-head bridging study for ES-SCLC in the U.S.
- Collaborated with **Syneos Health**, preparing for the prelaunch of Serplulimab injection (PD-1) in the U.S.

Med Tech: Sisram North American direct sales achieved revenue of **USD140 million (+28.2% YoY)**, accounting for approximately **40.5%** of Sisram's total revenue in 2022



Global Operation (2/2)



Leveraging global resources, quickly realizing and maximizing product value



Corporate Governance – Sustainable Development



Green growth and sustainable development

- Established EHS Committee to continuously improve EHS policies and set the 2nd EHS five-year strategic goals (2021-2026)
- Invested RMB1.15 million in special fund for water conservation in 2022, with a total annual water saving of $337,806 m^3$, 3.2% of the total annual water consumption

Improvement of product accessibility and affordability, taken the interest of stakeholders into consideration

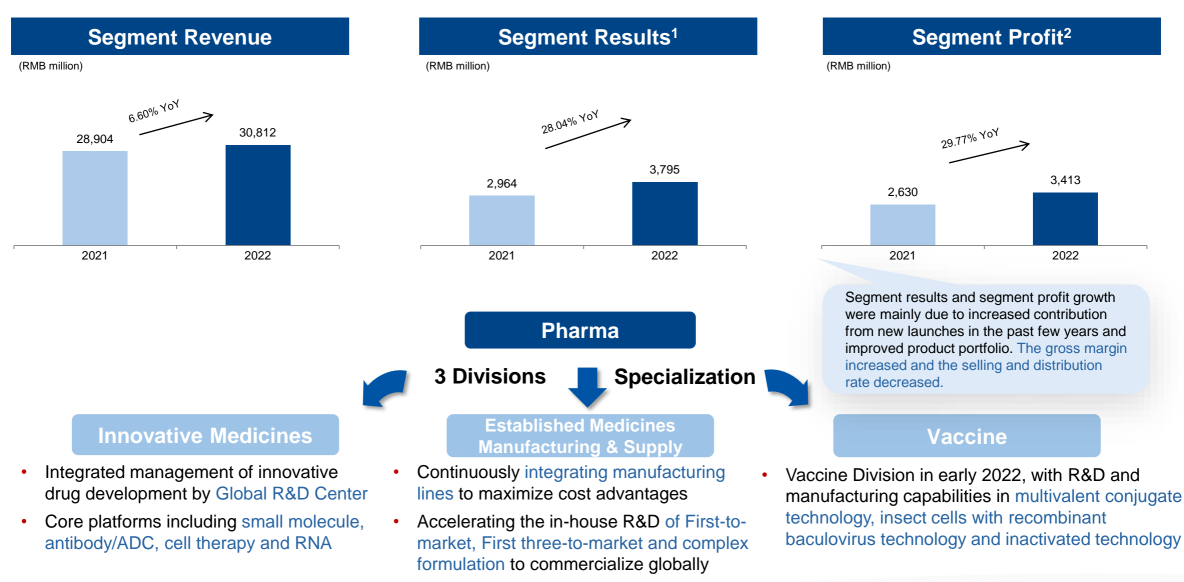
- Well-established systems for R&D, product quality management, staff training, social welfare and supply chain management
- Launched 2 orphan drugs/drugs for rare diseases, Aminohexanoic acid powder and Avatrobopag tablet; increased the accessibility of Ejilunsai injection (CAR-T) through commercial insurances and citizen insurances; in-house developed Antimalarial Series including Artesunate saved more than 56 million patients with severe malaria

Strengthen corporate governance with ESG to achieve sustainable development

- Established ESG Committee at the Board level; the independent Anti-Corruption Supervision Department (ACSD) designed a comprehensive anti-corruption system
- Published over 10 documents related to corporate governance on the official website
- Upheld the professional, branded, digital and compliant marketing system control

Pharmaceutical

Pharma - Performance



Note 1: segment results are obtained as segment revenue less costs of sales, selling and distribution expenses, administrative expenses and R&D expenses Note 2: pharmaceutical profit excludes the effect on sales of BNTX shares

Pharma - Core Product Revenue in Different Therapeutic Areas in 2022

Anti-tumor and Immune Modulation

RMB5,522 million 26%*

Revenue increase from Trastuzumab Injection (HER2), Avatrombopagmaleate Tablets, Adalimumab injection and from new launches in the past few years including Serplulimab Injection (PD-1) and Netupitant-Palonosetron

Anti-infection

RMB8,582 million 40%* (-0.45% YoY)

Mainly due to the combined effect of the decrease in the sales volume of Comirnaty (mRNA COVID-19 vaccine) and Micafungin, the revenue contribution from new products Azvudine tablets, Cravit (levofloxacin tablets and levofloxacin injection)

Metabolism and Alimentary System

RMB2,883 million 13%* (-0.24% YoY)

Mainly due to the impact of the execution of centralized procurement for Thioctic acid injection and Glutathione for injection

Cardiovascular System

RMB2,115 million 10%* (+6.12% YoY)

Mainly due to the increase in the sales volume of heparin series preparations

Central Nervous System

RMB1,003 million 5%* (-11.79% YoY)

Mainly due to the decline in sales volume of deproteinised calf blood serum injection

APIs and Intermediate Products

RMB1,248 million 6%* (+9.96% YoY)

Mainly due to the increase in the sales volume of amino acid series

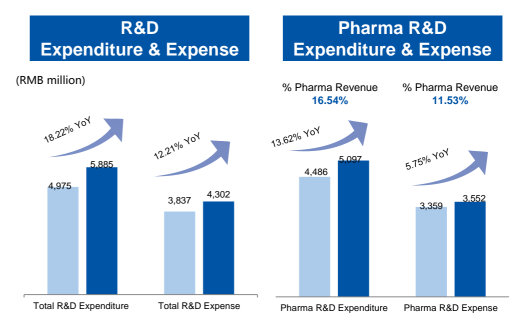


Note: represents core product revenue in single therapeutic area / sum of core product revenue in all therapeutic areas

Pharma - R&D Expenditure

R&D expenditure drives product portfolio optimization

- Phama R&D expenditure was RMB5,097 million (+13.62% YoY) in 2022, accounts for over 85% of the total R&D expenditure and 16.47% of the pharma revenue; Pharma R&D expense was RMB3,552 million, accounts for 11.53% of the pharma revenue
- new launches in the past few years including Serplulimab injection (PD-1), Trastuzumab injection (HER2), Avatrombopag tablets and Azvudine tablets accounts for over 30% of the pharma revenue, optimizing product portfolio
- Over 260 pipeline drugs in innovative drugs, biosimilars, generic drugs, consistency evaluation items, etc. by the end of 2022; received 249 applied pharma patents, including 16 U.S. patent applications, 17 PCT applications and 48 licensed invention patents in 2022



2021 2022



Pharma Key Progress - Serplulimab Injection

The first PD-1 inhibitor approved for first-line treatment of SCLC



1Q2023 Revenue

RMB250 million

2022 Revenue RMB340 million (Launched for 9 months)



Target: PD-1 Approved Indications in Chinese Mainland:

- MSI-H
- sqNSCLC
- ES-SCLC

Overseas Progress

- SCLC is granted with Orphan-drug Designation from FDA and EC
- Initiated ES-SCLC head-to-head bridging in the U.S.
- The MAA of ES-SCLC was accepted by the EMA

Outstanding Results

- Serplulimab + chemo (ES-SCLC) randomized, double-blind, median progression, global multi-center Phase 3 clinical data: Median OS 15.4 months, vs 10.9 month with placebo; 2 year OS rate 43.1%, vs 7.9% with placebo
- The clinical data have been published in world's top medical journals including The Journal of the American Medical Association (JAMA), Nature Medicine and British Journal of Cancer



Quick Market Access and Accelerated Market Penetration

- Completed tenders on procurement platforms in 27 provinces; covered 30% of the top 110 hospitals
- Commercialization team of about 400 people with experience in oncology drugs market
- Established efficient distribution network; maximized accessibility by leveraging DTP pharmacies and infusion centers



Pharma Key Progress – In-house R&D Vaccine Platform

Established Vaccine Subdivision in early 2022, with multivalent conjugate technology, insect cells with recombinant baculovirus technology and inactivated technology, to R&D and manufacture vaccines

Multivalent Conjugate Vaccine

Key progress

- Completed the enrollment of the Ph3 clinical trial of 13-valent pneumococcal conjugate vaccines (multivalent combinations) in Chinese Mainland in April 2023
- Fosun Adgenvax received Drug Manufacturing License from Sichuan Medical Products Administration in January 2023, laying the foundation for commercial manufacturing of vaccines under development

Received National Intellectual Property Rights

- The only multivalent combination technology with national patent and independent intellectual property rights
- 13-Valent Pneumococcal Conjugate
 Vaccine is the only pneumonia vaccine
 listed as a national major project





- Stable antigen structure, rapid immune response, earlier protection, complex antigen structure to enhance immunogenicity and reduce interference with different types of antigen immune response
- Cost and Safety Advantages
- Meat-free medium for carrier protein to reduce manufacturing cost and period

R&D Projects

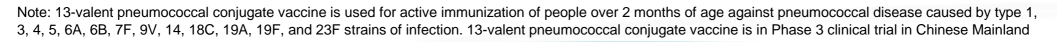
Inactivated Technology

- Launched Human Rabies Vaccine (Vero Cells) and Influenza Vaccine
- Nearly 20 years experiences in stable commercial manufacturing; innovative technology center for inactivated vaccines in Liaoning Province

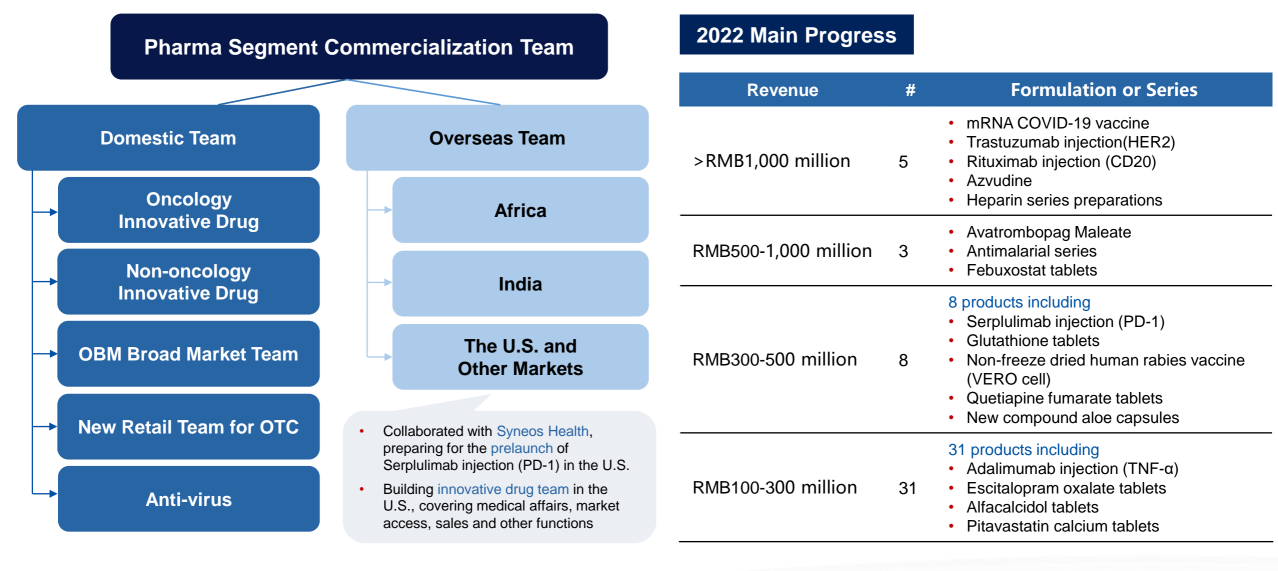
Insect Cells with Recombinant Baculovirus Technology

 First approved Hi5 insect cell line without nodavirus and SF9 insect cell line without rhabdoviruses in Chinese Mainland





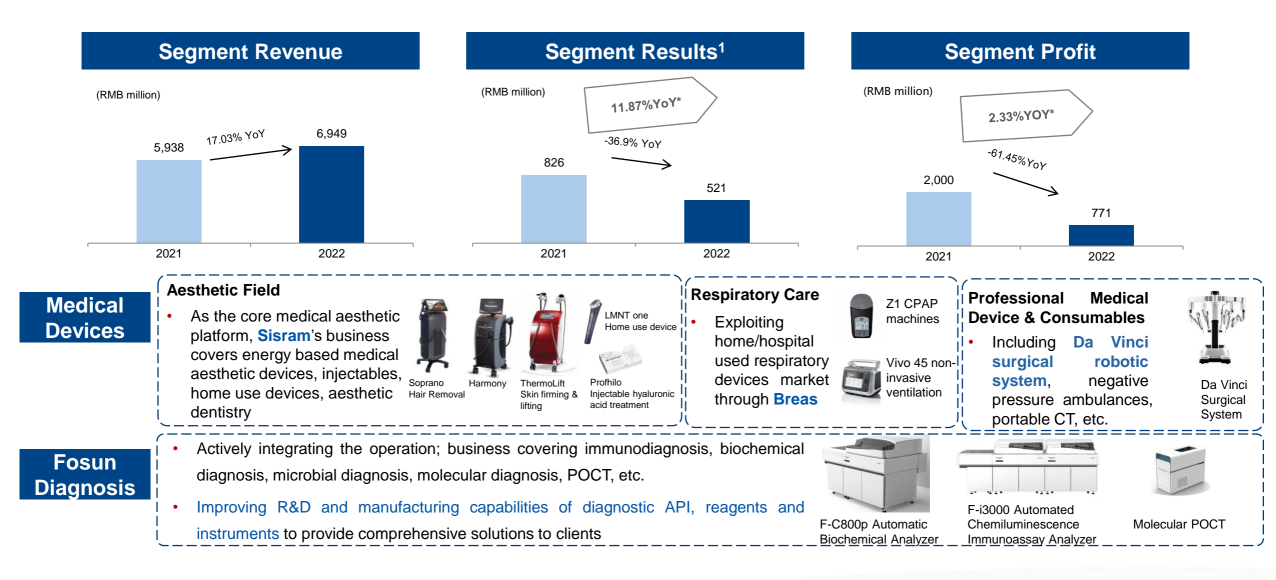
Pharma - Global Commercialization System





Med Tech

Med Tech - Performance





Note 1: Segment results are obtained as segment revenue less costs of sales, selling and distribution expenses, administrative expenses and R&D expenses Note 2: Segment results increased by 11.87% YoY, segment profit increased by 2.33% YoY, excluding the impact from equity transfer of Yaneng Bioscience

Medical Devices – Sisram Medical

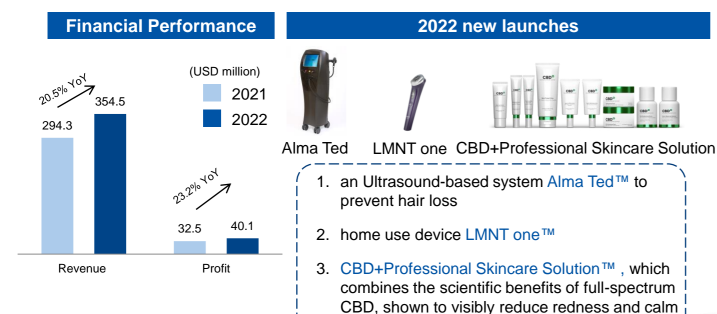
Establishing global Wellness Ecosystem based on energy-based devices and extending to injectables, aesthetic dentistry and personal care

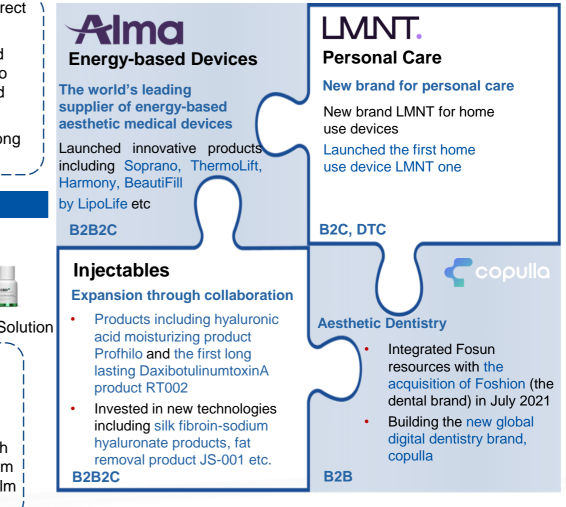
2022 Main Progress

Direct sales revenue accounts for 66% of the total revenue in 2022 (62% in 2021) with direct sales team covers 10 markets

- By entering into an agreement with PhotonMed and proposing to control the brand and channels of "PhotonMed" through M&A of assets, controlled subsidiary Sisram plans to build direct sales team for energy-based devices in **China** to further promote the brand
- Built new direct sales team in Dubai in February 2023 to develop and increase brand awareness in the Middle East; built new direct sales team in the UK to support the strong demand growth for product and services in Europe

the appearance of stressed skin







Medical Devices - Intuitive Fosun

Localization Process

	·	Made in China Joint R&D	•
Future		Localization in technology, manufacturing and services	•
2022	•	Building da Vinci Surgical Manufacturing R&D Center in Shanghai, covering about 31.2 acres	
2021	•	Da Vinci Innovation Center opened with 1,700 m ² of space to provide high-quality hands-on precision medicine training to approximately 4,000 doctors per year	•
2020	•	Da Vinci Surgical System test drives in more than 10 cities across China, with more than 800 doctors from nearly 200 hospitals participated in the experience	•
2019	+	Marketing the 4th generation Da Vinci XI Surgical System	
2017	ţ	Announced to form a JV with Intuitive Surgical in China in 2016 based on the long-term partnership and established Intuitive Fosun in Shanghai in 2017	

Global Commercialization

Main Products

Da Vinci Surgical System



- **55** da Vinci Surgical Systems were installed in China in 2022. By the end of 2022, **over 300 Systems** were installed in Chinese Mainland, Hong Kong and Macau regions and completed more than 100,000 surgeries within 2022
- As of June 30th 2022, 7,544 systems were installed worldwide, with more than 55,000 doctors trained to use the system, and performed over 10 million surgeries.

Ion Endoluminal System

- The robotic-assisted bronchoscopy platform, lon, was approved by FDA in 2019
- The lon guided lung nodule biopsy clinical feasibility trial completed enrollment at Shanghai Chest Hospital in October 2021. It is the first clinical trial using lon outside the United States





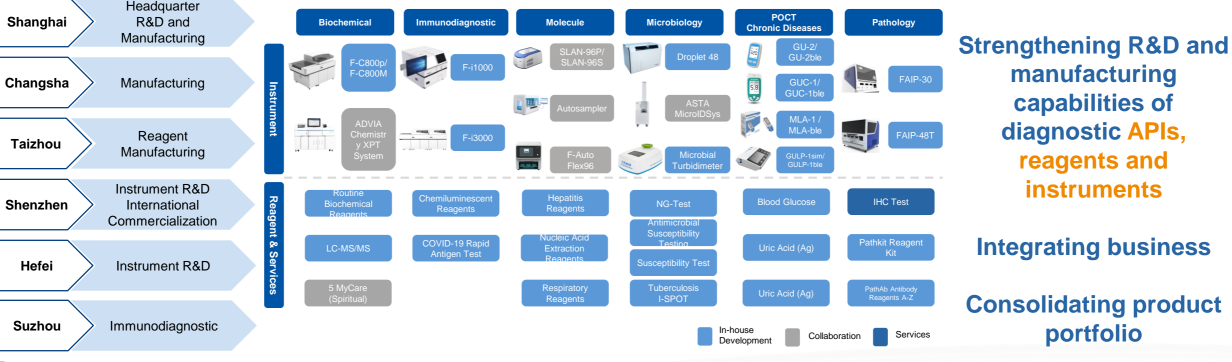
Medical Diagnosis - Core Products

Medical Diagnosis 2022 Major Progress

6 R&D and Manufacturing Bases

- Promoting the integration of medical diagnosis segment, constructing 6 R&D and manufacturing bases; R&D personnel account for more than 15% of the total number of Medical Diagnosis employees
- F-C800p Automatic Biochemical Analyzer launched in June 2022, together with the F-i3000 Automated Chemiluminescence Immunoassay Analyzer, formed Fosun Diagnostics biochemical immunoassay pipeline to meet the clinical diagnostic testing needs
- Self-developed COVID-19 Rapid Antigen Test was approved by NMPA in April 2022. It has received EU CE certification and has been included in the EU Common list of COVID-19 antigen tests and completed BfArM registration in Germany
- Self-developed Monkeypox PCR Detection Kit received EU CE certification in May 2022

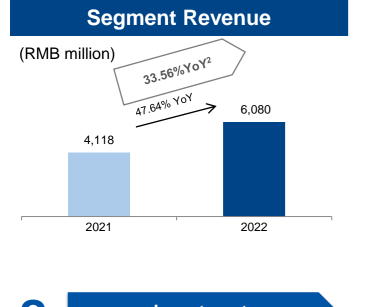
Instruments E APIs Reagents





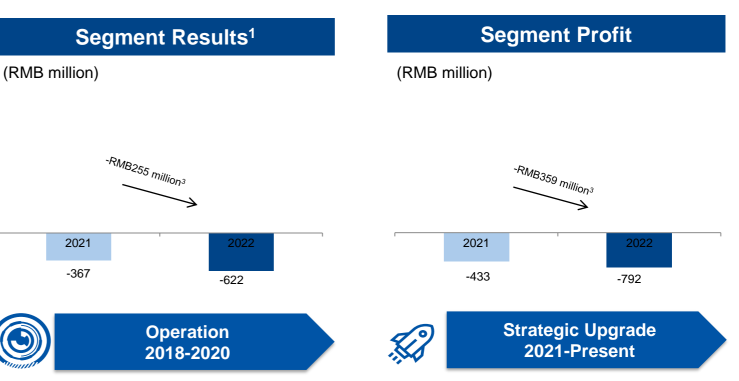
Healthcare Services

Healthcare Service - Performance



Investment 2011-2017

- Built offline healthcare network
- Gained experience in high-end healthcare
- Launched online healthcare services
- Developed regional medical centers



- Created advantageous specialty areas
- Online and offline strategic synergy
- Developed high-end aesthetic medical business
- Constructing specialties for health and wellness
- Integrating resources to build Internet healthcare ecosystem
- Consolidating the leading position as non-public healthcare provider
- Building intelligent Cloud Healthcare
- Building healthcare ecosystem

Note 1: Segment results are obtained as segment revenue less costs of sales, selling and distribution expenses, administrative expenses and R&D expenses Note 2: the revenue growth was mainly due to the growth of online business and the recovery of offline hospitals revenue. Segment revenue increased 33.56% YoY, excluding the impact of acquiring Guangzhou Xinshi Hospital

Note 3: the decrease of segment results and segment profit was mainly due to the investment in online business, periodic decrease in diagnosis and treatment volume of hospitals and initial loss of newly opened hospitals



Healthcare Services - Offline Services

Highlights

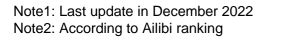


Covered Region

- Focus on the Yangtze River Delta, the Greater Bay Area and other regions; connecting medical centers with regional medical associations; integrating hospital resources
- 6,333 beds¹ in hospitals controlled by the Group by the end of 2022

Competitiveness

- Foshan Chancheng Hospital received JCI certification and ranked the TOP1 non-public hospital in China for 5 consecutive years²
- Shenzhen Hengsheng Hospital was granted JVF license



Pearl River Delta

Regional flagship hospitals include Foshan Chancheng, Shenzhen Hengsheng, etc.



JCI国际认证医院 Organization Accredited by Joint Commission International

Major Hospitals

- Class III General Hospital with 1,750 beds
- Realized revenue of RMB2,145 million, and profit of RMB111 million in 2022
- Fosun Pharma currently holds 86.47% of the share



- Class III General Hospital with 600 beds
- Acquired 60% stake of Shenzhen Hengsheng Hospital for RMB909 million in November 2017

- Class III General Hospital with 800 beds and over 900 doctors and employees
- Acquired 70% stake of Guangdong Xinshi Hospital in January 2022

Other Strategic Region









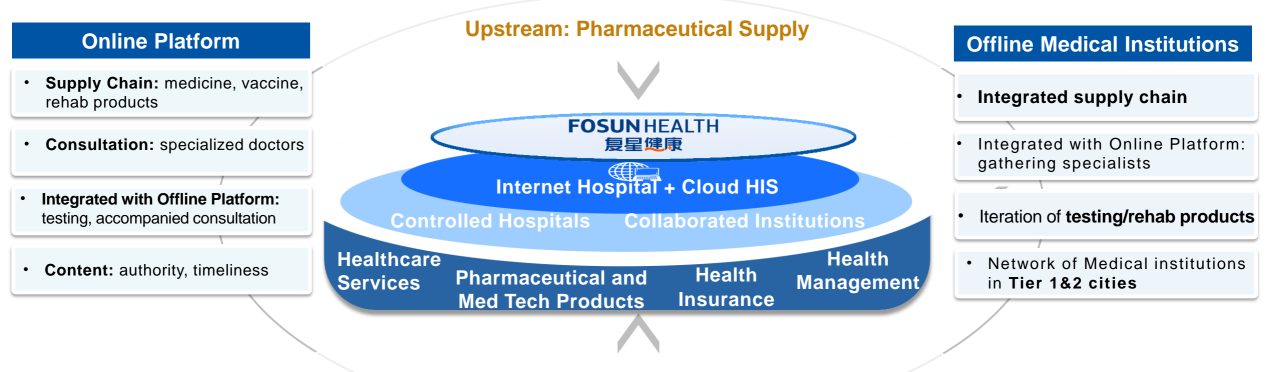




Healthcare Services – Integrating Online and Offline Services

- Integrated online and offline healthcare services from 2021, has received 10 internet hospital licenses as for now
- Building online medical service platform to provide healthcare services, pharmaceutical and med tech e-commerce, health insurance service and health management services

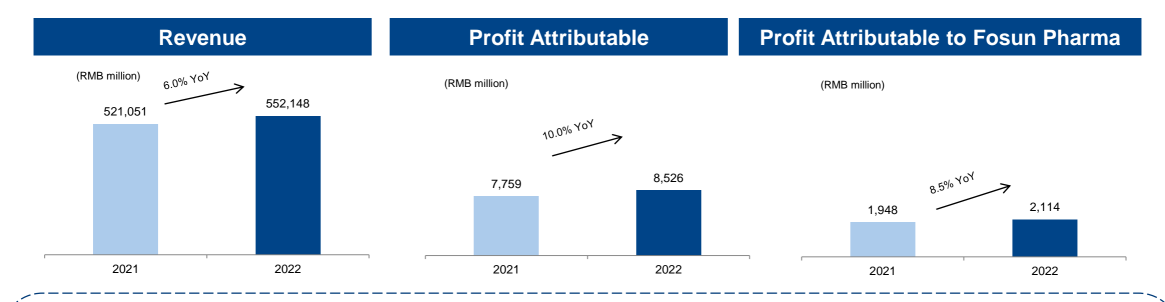
Accelerating online and offline services integration Building a one-stop healthcare management FHMO



Downstream: Insurance Payment



Sinopharm Performance



- Actively complied with the industry transformation trend, strengthened service capability of distribution network, and ensured the steady
 growth of key regions and markets while continuously improving the coverage and penetration ratio of business network. The 2022 revenue
 from the pharmaceutical distribution segment reached RMB406.60 billion (+4.27% YoY)
- Fully utilized advantages of "covering the whole country" logistics network, actively expanded derivative services while safeguarding
 personal protective products for COVID-19, and further enhanced the market share. The 2022 revenue from the medical device segment
 amounted to RMB120.85 billion (+11.77% YoY)
- Actively responded to the national strategy, undertook the new transformation and demand of separation of medical services and pharmaceutical sales, increased the allocation of resources, and made great efforts to promote the balanced development of professional pharmacies and traditional pharmacies. The 2022 revenue from retail pharmacy business reached RMB33.0 billion (+13.49% YoY)



Appendix

Large Molecules Pipeline (1/2)

Therapeutic Area	Product		Target/MOA Indication		Pre- Clinical	IND	Phase 1	Phase 2	Phase 3	NDA
				Squamous non-small cell lung cancer 1L	Global multi-center clinical trial Ph3, approved in Chinese Mainland in November 2022					
				Extensive-stage small cell lung cancer 1L	The MAA was accepted by the EMA; first U.S. bridging study subject had been dosed in November 2022; Granted Orphan-drug Designation by FDA and EC; approved in Chinese Mainland in January 2023					
		+Chemo	PD-1	Metastatic esophageal squamous-cell carcinoma 1L	-					
				Limited-stage small cell lung cancer	Global multi-center clinical trial Ph3; first subject had been dosed in the U.S. in January 2023 first subject had been dosed in Chinese Mainland in May 2022					
	HLX10 ¹			Neo-/adjuvant treatment of gastric cancer						
	(Serplulimab)	+Bevacizumab		Non-squamous non-small cell lung cancer 1L						
				Hepatocellular carcinoma 1L						
				Metastatic colorectal cancer 1L						
Anti-tumor		+HLX07	PD-1+EGFR	Squamous-cell carcinoma of the head and neck 2L						
			PD-1+EGFK	Squamous non-small cell lung cancer 1L	First subject had bee	n dosed in January	2022		•	
	HLX04-O ²		VEGF	Wet age-related macular degeneration			subject had been dos , Europe and Chinese		uary 2022;	
	HLX22 +Trastuzumab		HER2+HER2	Gastric cancer			nland in September 20		•	
	HLX07		EGFR	Solid tumors (non-small cell lung cancer, esophageal carcinoma, etc.)	Approved clinical tria	ls by FDA			•	
	HLX11 (Pertuz	zumab) ³	HER2	Breast cancer	Global multi-center cl	nical trial Ph3; first	subject had been dos	ed in Chinese Mainla	nd in 2022	
	HLX05 (Cetuximab) ⁴		EGFR	Metastatic colorectal cancer and squamous-cell carcinoma of the head and neck						
	HLX02 (Trastuzumab)		HER2	Breast cancer and metastatic gastric cancer	The BLA was accepte approved in Europe a		nd in 2020			

Note 1: granted KG Bio to develop and commercialize HLX10 in 10 countries in Southeast Asia

Note 2: granted ESSEX an exclusive license to develop, manufacture, and commercialize HLX04 in human ophthalmic therapeutic use

Note 3: granted Organon exclusive global commercialization rights except for China

Note 4: granted Jingze Biotech to commercialize HLX05 in China

Note 5: collaborated with Accord, Cipla, Jacobson, mAbxience, Eurofarma and Abbott

Note 6: last update on 28th April 2023



Large Molecules Pipeline (2/2)

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Therapeutic Area	Product	Target/MOA	Indication	Pre- Clinical	IND	Phase 1	Phase 2	Phase 3	NDA
	50 (500		HER2-positive advanced malignant solid tumor						
	FS-1502	HER2	HER2-positive locally advanced or metastatic breast cancer						
	FS-1502+Serplulimab	HER2+PD-1	Advanced gastric cancer with HER2 expression						
	HLX14 (Denosumab) ¹	RANKL	Osteoporosis	Initiated Ph3 clinica	al trial in Chinese N	lainland in June 202	2; approved to enter	Ph3 clinical trial by	TGA in July 2022
Anti-tumor	HLX26	LAG-3	Solid tumors and lymphomas						
	HLX35 ²	EGFR×4-1BB	Solid tumors	Approved to enter	clinical trials by NN	/IPA in January 2022	; first subject had be	en dosed in Chines	e Mainland in June 2
	HLX301	PD-L1×TIGIT	Solid tumors			ralia in February 202 IPA in March 2022; f		n dosed in Chinese	Mainland in July 202
	HLX15 (Daratumumab)	CD38	Multiple myeloma	First subject had b	een dosed in Chine	ese Mainland in Febr	ruary 2023		
	HLX13 (Ipilimumab)	CTLA-4	Melanoma, renal cell carcinoma and metastatic colorectal cancer				-		
Blood system	Recombinant Human Erythropoietin Injection (pre-filled syringe)	EPO	Anemia of renal disease						
	Recombinant Insulin Glargine Injection	INSR	Diabetes						
Metabolism and Digestive System	Mixed Protamine Zinc Recombinant Insulin Lispro Injection (50R)	INSR	Diabetes						
	Liraglutide Injection	GLP-1	Diabetes						
Others	RT002	Bio 1 Bio 1	Moderate to severe glabellar lines in adults (GL) Cervical dystonia (CD)			in April 2023 in Chinese Mainland	d in January 2022		-

Note 1: granted Organon exclusive global commercialization rights except for China Note 3: last update on 28th April 2023 Note 2: granted Binacea to research, develop, manufacture and commercialize the HLX35 globally except for China (including Hong Kong, Macau and Taiwan region)

Small Molecules Pipeline (1/2)

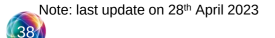
Therapeutic Area	Project	Target/MO A	Indication	Pre-Clinical	IND	Phase 1	Phase 2	Phase 3	NDA	
			Breast cancer (1L)	Approved to enter Ph3 clinical trial by NMPA in January 2022; Ph1 clinical trial in the U.S.						
	FCN-437c	CDK4/6	Breast cancer (2L)	Approved to enter Ph3 of	clinical trial by NMPA	in January 2022; appro	ved to enter clinical trial	s by FDA	•	
	SAF-189	ALK	Non-small cell lung cancer	Initiated Ph3 clinical trial in Chinese Mainland in January 2022; Ph1 clinical trial in the U.S.						
	SAF-109	ROS1	Non-small cell lung cancer	Approved to enter clinica						
	HLX-208	BRAF V600E	Solid tumors (metastatic colorectal cancer, non- small cell lung cancer, etc.) LCH and ECD5	Granted with the Breakt Approved to enter Ph1b	hrough Therapy Desi /Ph2 clinical trials by	gnation by the NMPA in NMPA in January 2022	April 2023;			
			eurofibromatosis type 1 Global multi-center clinical trial							
		I-159 MEK	Low-grade glioma							
Anti-tumor			Arteriovenous malformation	Approved to enter clinic	al trials by NMPA in N	/lay 2022				
Anti-tumor	FCIN-139		Histiocytic tumor	Granted with the Breakthrough Therapy Designation by the NMPA in April 2023;						
			Langerhans cell histiocytosis	Approved to enter clinical trials by NMPA in May 2022 Approved to enter Ph2 clinical trial by NMPA in March 2023						
			Langerhans cell histiocytosis in children	Approved to enter Ph2 of		•				
	YP01001	VEGFR	Advanced solid tumor							
			Myeloid malignancy	Approved to enter Ph2 of	clinical trials by NMPA	in March 2023				
	FCN-338	-338 BCL-2	Hematological malignancy	Approved to enter Ph1 clinical trial in the U.S.						
			Relapsed or refractory B-cell lymphoma				•			
	FH-2001	FGFR/PD- L1	Advanced malignant solid tumors							

Note: last update on 28th April 2023

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Small Molecules Pipeline (2/2)

Therapeutic Area	Project	Target/MOA	Indication	Pre-Clinical	IND	Phase 1	Phase 2	Phase 3	NDA		
Pland System	Avatrombopag Tablet	TPO-R	Chronic idiopathic thrombocytopenic purpura	NDA was accepted by NMPA in December 2022							
Blood System	Tenapanor Tablet	NHE 3	End-stage Renal Disease – Hemodialysis								
Metabolism	Ferric Pyrophosphate Citrate	-	Iron replacement for HD patients								
and Digestive System	Tenapanor Tablet	NHE 3	Irritable Bowel Syndrome with Constipation	Ph1 clinical trial in Chinese Mainland; NDA in Hong Kong and Macau regions							
-,	FCN-342	URAT1	Gout								
	Molnupiravir	RNA polymerase	Treatment of COVID-19								
	Paxlovid	3CL Protease	Treatment of COVID-19								
Infectious Diseases	mRNA COVID-19 BNT162b2 & bivalent vaccine	-	Immunization to prevent COVID-19	Administrated in Hong	g Kong, Macau and	Taiwan regions		•			
	PA-824	-	XDR – Tuberculosis MDR – Tuberculosis	Launched Pretomanic	I in the U.S.*						
Nervous System	Opicapone Tablet	COMT	Parkinson's syndromes	Launched Ongentys i	n Europe*			_			
Others	Fortacin spray (Lidocaine Prilocaine spray)	-	Premature ejaculation	Launched in Europe*							
	ET-26	-	Anesthesia	Approved to enter Ph	2 clinical trial by NN	IPA in July 2022					



Vaccine Pipeline

Product	Technology	Pre-Clinical	IND	Phase 1	Phase 2	Phase 3	NDA
Freeze-dried Human Rabies Vaccine (Vero Cells)	Inactivated						
4-Valent Influenza Vaccine	Inactivated						
Human Diploid Cell Rabies Vaccine	Inactivated						
13-Valent Pneumococcal Conjugate Vaccine	Multivalent Conjugate						
24-Valent Pneumococcal Conjugate Vaccine	Multivalent Conjugate						
23-Valent Pneumococcal Conjugate Vaccine	Multivalent Conjugate						
Quadrivalent Meningococcal Polysaccharide Vaccine	Multivalent Conjugate						
Tetanus Vaccine	-						
Quadravalent Meningococcal Conjugate Vaccine	Multivalent Conjugate						
Recombinant Zoster Vaccine	Insect Cells with Recombinant Baculovirus						
Recombinant Quadravalent Influenza Vaccine	Insect Cells with Recombinant Baculovirus						

Note: last update on 28th April 2023





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