



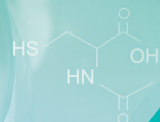
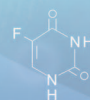
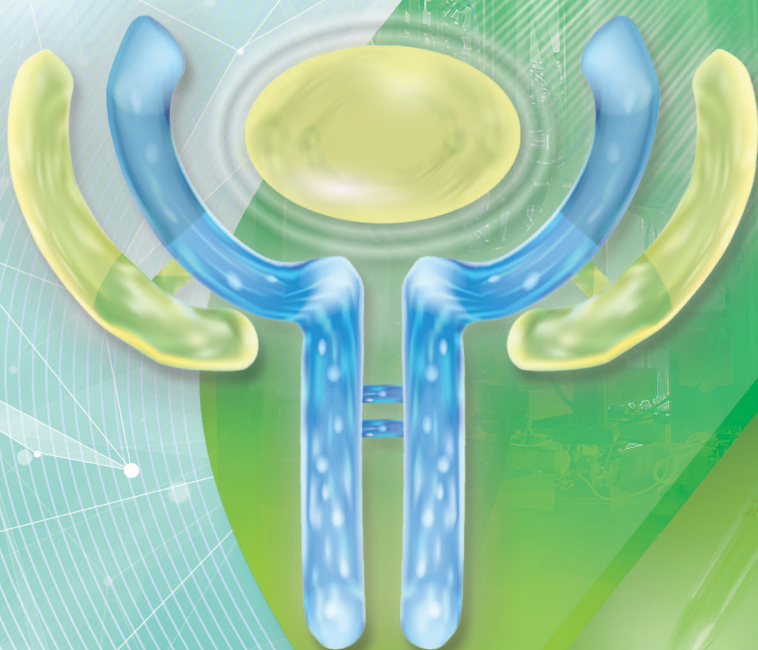
Sunho Biologics, Inc.

盛禾生物控股有限公司

(Incorporated in the Cayman Islands with limited liability)

Stock Code : 2898

INTERIM REPORT
2025



CONTENTS

2	Corporate Information
4	Management Discussion and Analysis
20	Other Information
25	Condensed Consolidated Statements of Profit or Loss and Other Comprehensive Income
26	Condensed Consolidated Statements of Financial Position
28	Condensed Consolidated Statements of Changes in Equity
29	Condensed Consolidated Statements of Cash Flows
30	Notes to the Condensed Consolidated Financial Statements
42	Definitions



Corporate Information

Company Name

Sunho Biologics, Inc. (盛禾生物控股有限公司)

Directors

Executive Directors

Mr. ZHANG Feng (張峰)

(Chairman)

Dr. YIN Liusong (殷劉松)

(Chief executive officer and chief scientific officer)

Ms. JIANG Xiaoling (姜曉玲)

(Vice president)

Non-executive Director

Mr. FAN Rongkui (范融奎)

Independent Non-executive Directors

Mr. CHAN Heung Wing Anthony (陳向榮)

Ms. FENG Lan (馮嵐)

Mr. SHI Luwen (史錄文)

Audit Committee

Mr. CHAN Heung Wing Anthony (陳向榮)

(Chairman)

Ms. FENG Lan (馮嵐)

Mr. SHI Luwen (史錄文)

Remuneration Committee

Ms. FENG Lan (馮嵐)

(Chairlady)

Mr. ZHANG Feng (張峰)

Mr. SHI Luwen (史錄文)

Nomination Committee

Mr. ZHANG Feng (張峰)

(Chairman)

Ms. FENG Lan (馮嵐)

Mr. SHI Luwen (史錄文)

Joint Company Secretaries

Ms. XU Chunqin (徐春芹)

Ms. WONG Hoi Ting (黃凱婷)

Authorised Representatives

Mr. ZHANG Feng (張峰)

Ms. WONG Hoi Ting (黃凱婷)

Auditor

ZHONGHUI ANDA CPA Limited (appointed with effect from September 16, 2025)

Certified Public Accountants

Registered Public Interest Entity Auditor

23/F, Tower 2

Enterprise Square Five

38 Wang Chiu Road

Kowloon Bay

Hong Kong

Legal Advisers

As to Hong Kong laws

**Eric Chow & Co. in Association with
Commerce & Finance Law Offices**

3401, Alexandra House

18 Chater Road

Central

Hong Kong

Registered Office in Cayman Islands

PO Box 309

Ugland House

Grand Cayman, KY1-1104

Cayman Islands

Headquarters

Room 302, Building 3

No. 198 Peninsula Middle Road, Dipu Street

Anji County, Huzhou City

Zhejiang Province

PRC

No. 5 Xingjian Road

Nanjing Economic and Technological

Development Zone

PRC

Corporate Information

Principal Place of Business in the PRC

No. 5 Xingjian Road
Nanjing Economic and Technological
Development Zone
PRC

Principal Place of Business in Hong Kong

31/F, Tower Two
Times Square
1 Matheson Street
Causeway Bay
Hong Kong

Principal Share Registrar and Transfer Office

Maples Fund Services (Cayman) Limited
PO Box 1093, Boundary Hall
Cricket Square
Grand Cayman KY1-1102
Cayman Islands

Hong Kong Share Registrar

Computershare Hong Kong Investor Services Limited
Shops 1712-1716, 17th Floor
Hopewell Centre
183 Queen's Road East, Wanchai
Hong Kong

Principal Bankers

**Bank of Communications Co., Ltd.,
Nanjing Xingang Development Zone Branch**
No. 10 Xingzhi Road
Qixia District, Nanjing City
Jiangsu Province
PRC

**China Merchants Bank Co., Ltd.,
Zhujiajiang Road Sub-branch**
No. 280 Zhujiajiang Road
Xuanwu District, Nanjing City
Jiangsu Province
PRC

Stock Code

2898

Company Website

www.sunho-bio.com.cn

Management Discussion and Analysis

Business Review

Founded in 2018, we are a clinical stage biopharmaceutical company that focuses on the discovery, development and commercialization of biologics for the treatment of cancers and autoimmune diseases. We have three Core Products, namely, IAH0968, IAP0971 and IAE0972, all of which are developed in-house. IAH0968 is an antibody-dependent cell mediated cytotoxicity ("**ADCC**") enhanced monoclonal antibody ("**mAb(s)**"), and we have initiated Phase II clinical trials for colorectal cancer ("**CRC**") and gastric cancer ("**GC**"). IAP0971 and IAE0972 are both immunocytokines and we have completed Phase I clinical trials for advanced solid tumors including non-small cell lung cancer ("**NSCLC**") and CRC. We have initiated Phase II clinical trials of IAE0972 for head and neck squamous cell carcinoma ("**HNSCC**") and nasopharyngeal carcinoma ("**NPC**").

R&D of product candidates

Our R&D capabilities cover development of candidates in the forms of mAbs, bispecific antibodies ("**bsAb(s)**"), and fusion proteins, some of which extend indications into treatment areas beyond oncology. Our Core Product IAH0968 is an ADCC enhanced mAb targeting human epidermal growth factor receptor 2 ("**HER2**") with 100% fucose knock out, which greatly enhances the binding affinity of its fragment crystallizable ("**Fc**") to its receptor FcγRIIIa. ADCC is an immune mechanism through which Fc receptor-bearing effector cells including natural killer ("**NK**") cells and CD+8 T cells can recognize and kill antibody-coated target cells expressing tumor- or pathogen-derived antigens on their surface. It is one of the most important methods for antibody drugs to kill tumor cells. The typical ADCC involves activation of NK cells by antibodies in a multi-tiered progression of immune control. A NK cell expresses Fcγ receptors ("**FcγR**"). These receptors recognize and bind to the Fc domain of an antibody, and the antigen binding fragment ("**Fab**") domain of which binds to the tumor associated antigen ("**TAA**") on the tumor cell. When both TAA and FcγR are engaged respectively by the Fab and Fc portions of the antibody, ADCC is initiated, since this creates a bridge from the tumor cell to the effector cell. However, the natural affinity between antibodies and FcγR is relatively weak, and Fc engineering to enhance affinity has become a common method.

Our featured products, immunocytokines, are designed through our proprietary and internally developed Armed ImmunoCytokine Platform ("**AIC™ Platform**") by our core R&D team in researching antibody-cytokine fusion proteins. They function through diverse mechanisms of action yet share a similar structure comprising an antibody or quasi-antibody moiety that targets tumors and blocks signaling pathways regulating tumor growth and proliferation, and cytokine payloads that activate the immune system within the tumor microenvironment ("**TME**"). Such a design is expected to overcome drawbacks of conventional cytokine-based drugs, such as short half-lives, systemic cytotoxicity and modest efficacy due to cytokine pleiotropy and off-target effects. It is expected to achieve enhanced antitumor effects through the synergy between the antibody and cytokine payloads, which potentially address the needs of cancer patients who suffer from disease progression related to the immunosuppressive TME and drug resistance.

Management Discussion and Analysis

IAH0968

Our Core Product IAH0968 is an internally developed, the first anti-HER2 antibody in clinical stage with 100% fucose-removal. Antibodies consist of two structural regions, Fab and Fc. Unlike the Fab region, which defines the specific target of an antibody, Fc region mediates ADCC by activating the immune system through engaging various Fc receptors. Studies of the structure of the Fc region of antibodies and its receptor FcγRIIIa complex revealed that the core fucose of the Fc region is accommodated at a place that interferes with the binding between the Fc region and FcγRIIIa, and thus reducing the affinity between them and resulting in lower ADCC activity. Therefore, modifying to remove fucose is desirable to better recruit immune cells, resulting in enhanced ADCC activity. As a result, this approach has been widely attempted in the biopharmaceutical industry. However, despite numerous attempts by multiple players to modify antibodies through various approaches, such as Fc point specific mutation and fucose removal, most resulting antibodies still contain a certain percentage of core fucose.

The Phase I clinical trial showed that IAH0968 was well tolerated and exhibited antitumor activities in patients with advanced HER2+ malignant solid tumors including breast cancers, gastric cancers, CRC and biliary tract carcinoma ("**BTC**") with drug resistance to trastuzumab, pertuzumab, cetuximab, docetaxel, oxaliplatin, capecitabine, irinotecan, nab-paclitaxel and apatinib, or anti-PD-1 mAbs. Data showed that only one dose-limiting toxicity ("**DLT**") was found at dosage 10mg/kg, and no maximum tolerable dose ("**MTD**") was reached. While no head-to-head study was conducted, the Phase I clinical data showed that IAH0968 achieved significantly improved objective response rate ("**ORR**") and disease control rate ("**DCR**") in heavily pretreated metastatic CRC and BTC patients, when compared to the historical data of current treatments. For heavily pretreated metastatic CRC and BTC patients, the ORR was 40%, and DCR was 80%.

We obtained the IND approval for conducting Phase I and Phase II clinical trials of IAH0968 from the NMPA in October 2020, commenced the Phase I clinical trial in August 2021, and completed the Phase I clinical trial of using IAH0968 as a monotherapy for heavily pretreated patients with advanced HER2+ malignant solid tumors in March 2023. Based on the encouraging clinical data from the Phase I trial, we obtained IND approvals from the NMPA to conduct Phase II and Phase III clinical trials of using IAH0968 in combination with chemotherapy for first-line treatment of inoperable HER2+ advanced or metastatic CRC, and to conduct Phase II clinical trials of using IAH0968 in combination with chemotherapy for first-line treatment of HER2+ metastatic BTC patients in September 2022. We also obtained IND approval from the NMPA to conduct Phase II and Phase III clinical trials of using IAH0968 in combination with chemotherapy for first-line treatment of HER2+ advanced or metastatic GC, and HER2-expressing solid tumors in April 2024. We have dosed the first CRC patient of the Phase IIa trial in May 2023. We entered a Phase IIb/III clinical trial for CRC in January 2024. We also entered a Phase IIb/III clinical trial for GC in August 2024. We also obtained IND approval from the NMPA to conduct Phase II and Phase III clinical trials of using IAH0968 in combination with chemotherapy for third-line treatment of HER2+ advanced CRC in April 2025.

Management Discussion and Analysis

IAP0971

Our Core Product IAP0971 is an internally developed, dual-moiety, anti-programmed death-1 ("**PD-1**") antibody-IL-15/IL-15R α heterodimer dual T cell and NK cell agonist. IAP0971 is expected to synergistically strengthen the antitumor activity through blockade of the PD-1/its ligand ("**PD-L1**") signaling pathway and accumulating IL-15 at the targeted tumor site to activate its nearby immune cells, including CD8+ T cells and NK cells, directly activating both innate and adaptive immune systems.

In July 2023, we completed Phase I clinical trial of IAP0971 for advanced malignant tumors. Phase I clinical data showed that IAP0971 exhibited a favorable safety profile at up to 200 μ g/kg in patients with advanced malignant tumors, with no DLT and MTD observed. Preliminary antitumor efficacy was observed in five patients treated with IAP0971 as later-line therapy. These five patients include one with CRC, one with cervical cancer, one with ovarian cancer, and two with NSCLC, and those patients underwent multiple rounds of treatments including chemotherapy, targeted therapy, immunotherapy and/or their combination, and experienced disease progress and metastases. After receiving IAP0971 for two treatment cycles, all five patients achieved stable disease ("**SD**"). Especially, one NSCLC patient complicated with adrenal gland and other metastases was resistant to several prior treatments, including chemotherapy regimes such as multiple paclitaxel-containing combination, and combination therapies with targeted therapy and immunotherapy, such as erlotinib, camrelizumab, sintilizumab and bevacizumab. This patient received 120 μ g/kg IAP0971 for two treatment cycles and achieved SD. The other NSCLC patient complicated with pleura or pleural effusion metastases was resistant to several prior treatments, and also achieved SD after two cycles of 200 μ g/kg IAP0971 administration.

In January 2022 and December 2021, we obtained IND approvals from both the NMPA and the FDA for conducting Phase I and Phase II clinical trials in patients with advanced malignant tumors, respectively. We commenced the Phase I clinical trial in China in June 2022 according to a protocol approved by both the NMPA and the FDA, and completed the Phase I clinical trial in July 2023. In May 2023 and August 2023, we also obtained IND approvals from both the NMPA and the FDA for conducting Phase I and Phase II clinical trials in patients with BCG unresponsive high risk non-muscle invasive bladder cancer ("**NMIBC**"), respectively. We dosed the first NMIBC patient in March 2024.

Management Discussion and Analysis

IAE0972

Our Core Product IAE0972 is an internally developed, dual-moiety, anti-epidermal growth factor receptor ("EGFR") antibody-IL-10 homodimer bifunctional fusion protein for immune cell activation. Like IAP0971, IAE0972 is also expected to achieve synergistical antitumor activities leveraging the advantages of immunocytokine yet through a different combination of antibody target and cytokine payload. It is designed to blockade the EGFR signaling pathway and specifically deliver IL-10 to the targeted tumor site to activate CD8+ T cells, and potentially NK cells.

We obtained the IND approval for conducting Phase I and Phase II clinical trials in patients with advanced solid tumors from the FDA and the NMPA in December 2021 and January 2022, respectively, commenced the Phase I clinical trial in China in June 2022 according to a protocol approved by both the NMPA and the FDA, and completed the Phase I clinical trial in July 2023. In our Phase I clinical trial of IAE0972 for advanced solid tumors, we recruited 14 patients with advanced esophageal squamous cell carcinoma, rectal cancer, gastric cancer, pancreatic cancer, small cell lung cancer ("SCLC") or NSCLC who progressed from at least one line of treatment. We completed dose escalation for 1µg/kg, 10µg/kg, 100µg/kg, 0.3mg/kg, 1.0mg/kg and 2.5mg/kg of IAE0972, and only observed one Grade 3 adverse events. No DLT occurred and MTD was not reached. Preliminary efficacy was observed in multiple heavily pretreated patients who failed all previous therapies. A CRC patient complicated by lung metastasis, who has received multiple lines of prior treatments including standard mFOLFOX6 (5-fluorouracil, leucovorin and oxaliplatin) and CapeOX (capecitabine and oxaliplatin) regimens, achieved SD after given 10µg/kg of IAE0972 for two treatment cycles. Another patient with rectal cancer and lung metastasis and lymph node metastasis, who had experienced recurrence after received two resections, achieved SD after receiving 1.0mg/kg of IAE0972 monotherapy for two cycles.

We also obtained the IND approval for conducting Phase II and Phase III clinical trials of IAE0972 in combination with chemotherapy in recurrent or metastatic HNSCC and NPC from the NMPA in September 2024. We have dosed the first patient of the Phase II clinical trial of HNSCC and NPC in May 2025.

Other pipeline products

In addition to our product candidates mentioned above, we are developing a number of clinical stage and IND-enabling product candidates that we believe have high commercial viability. As of June 30, 2025, except for IBC0966, we maintained the global rights to develop and commercialize them. For IBC0966, we have exclusive rights to develop, manufacture and commercialize in Greater China including Mainland China, Hong Kong, Macau and Taiwan and have partial overseas rights.

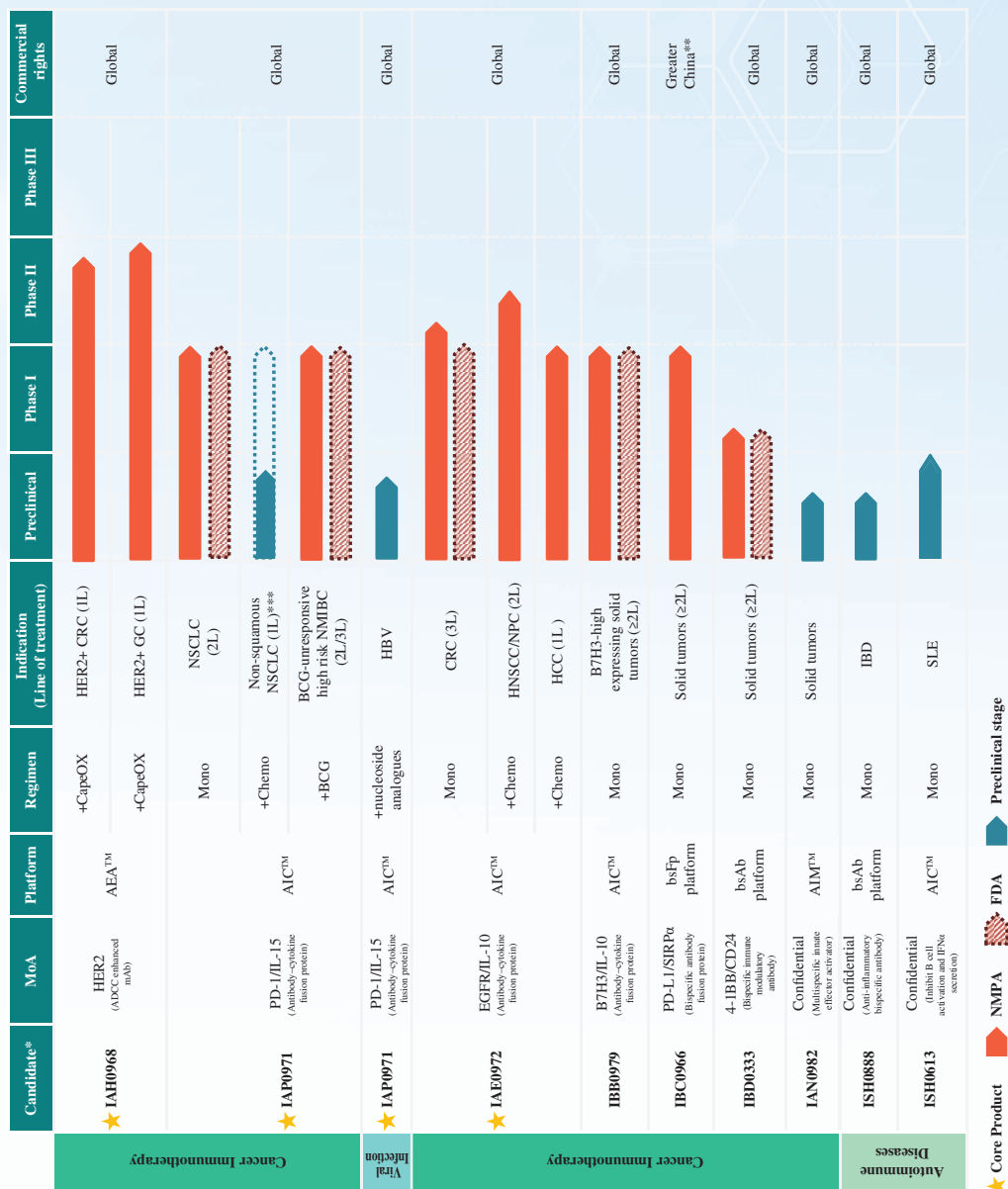
- **IBB0979:** IBB0979, another immunocytokine developed by us, is a clinical stage, dual-moiety, anti-B7H3 antibody-IL-10 homodimer bifunctional fusion protein for immune cell activation. It is designed to bind to B7H3 and trigger blockage of downstream signaling pathways that participate in TME shaping and development, and deliver IL-10 to activate CD8+ T cells to fight against tumors. We obtained the approval for conducting Phase I and Phase II clinical trials in patients with locally-advanced or metastatic solid tumors from the FDA and the NMPA in October 2022 and November 2022, respectively. The Phase I clinical trial is currently on-going, with the first patient dosed in July 2023. Since B7H3 is overexpressed in a wide range of cancers including glioma, thyroid, lung, head and neck, rectal, prostate, breast, skin, renal cell, and ovarian cancers, it has the potential to become a next-generation therapy for resolving T cell exhaustion in cancer patients. We have obtained IND approval from the NMPA for conducting Phase II and Phase III clinical trials of IBB0979 in combination with chemotherapy in recurrent or metastatic SCLC in April 2025.

Management Discussion and Analysis

- **IBC0966:** IBC0966 is a clinical stage anti-PD-L1 antibody-SIRP α bifunctional fusion protein that simultaneously stimulates both innate and adaptive immunity to achieve strong synergistic effects and induce long-lasting tumor-specific immune responses. It is designed to bind to PD-L1 and trigger blockage of the PD-1/PD-L1 signaling pathway to enable T cells to recognize and kill targeted cancer cells, and in the meantime deliver SIRP α to the targeted TME to interact with CD47 to block the “don’t eat me” signal of macrophages for tumor cell killing. In March 2021, we obtained the IND approval from the NMPA for conducting clinical trials of IBC0966. We completed the Phase I clinical trial of IBC0966 as monotherapy for advanced malignant tumors in December 2023. We acquired exclusive rights from ImmuneOnco Biopharmaceuticals (Shanghai) Inc. (“ImmuneOnco”) to develop, manufacture and commercialize IBC0966 in Greater China including Mainland China, Hong Kong, Macau and Taiwan.
- **IBD0333:** IBD0333 is a clinical stage 4-1BB and CD24 bsAb that simultaneously stimulates both innate and adaptive immunity to achieve strong synergistic effects with reduced hepatotoxicity. It is designed to bind to 4-1BB, a robust immune cell activator expressed by CD8+ T cells as well as DC cells, monocytes, B cells, mast cells, NK cells and neutrophils, and CD24, a promising target that plays a key role in tumor evasion in CD24-sialic-acid-binding Ig-like lectin 10 axis and thus is highly expressed in many cancer types. We have obtained IND approvals from the FDA in June 2023 and from the NMPA in July 2023. We initiated a Phase I clinical study in March 2024 in patients with locally advanced/metastatic solid tumors.
- **ISH0613:** ISH0613 is an internally developed bifunctional antibody fusion protein that simultaneously inhibits B cell activation and IFN α secretion based on our AICTM Platform. We are developing ISH0613 as a monotherapy for the treatment of systemic lupus erythematosus (“SLE”). We have filed IND application to the FDA for conducting Phase I clinical trial of ISH0613 in June 2025.
- **IAN0982:** IAN0982 is an internally developed multi-specific innate effector activator based on our AIMTM Platform. We are developing IAN0982 as a monotherapy or in combination with other therapeutics including chemotherapy and immunotherapy for the treatment of advanced solid tumors.
- **ISH0888:** ISH0888 is an internally developed bifunctional anti-inflammatory bispecific antibody based on our bsAb Platform. We are developing ISH0888 as a monotherapy for the treatment of inflammatory bowel disease.

Management Discussion and Analysis

The following diagram summarizes the status of the product pipeline of the Group as of June 30, 2025:



Management Discussion and Analysis

Abbreviations: 1L = first-line; 2L = second-line; 3L = third-line; ADCC = antibody-dependent cell-mediated cytotoxicity; AEA™ = ADCC Enhanced Antibody Platform; AIC™ = Armed ImmunoCytokine Platform; AIM™ = Armed Innate Effector Multi-specific Platform; BCG = Bacillus Calmette-Guerin; bsAb = bispecific antibody; bsFp = bispecific fusion protein; CapeOX = capecitabine and oxaliplatin; Chemo = chemotherapy; FDA = U.S. Food and Drug Administration; GC = gastric cancer; mAb = monoclonal antibody; Mono = monotherapy; NMPA = National Medical Products Administration; NSCLC = non-small cell lung cancer; NMIBC = non-muscle invasive bladder cancer; CRC = colorectal cancer; HBV = hepatitis B virus; HNSCC = head and neck squamous cell carcinoma; NPC = nasopharyngeal carcinoma; HCC = hepatocellular carcinoma; IBD = inflammatory bowel disease; SLE = systemic lupus erythematosus.

Notes:

- * All the product candidates are administered intravenously, except for IAP0971 for the treatment of 2L/3L NMIBC, which will be administered through intravesical instillation, as well as IAP0971 for the treatment of NSCLC, which will be administered through subcutaneous injection.
- ** We acquired exclusive rights from ImmuneOnco to develop, manufacture and commercialize IBC0966 in Greater China including Mainland China, Hong Kong, Macau and Taiwan, as well as 7.5% of interests in the overseas rights of IBC0966. For more information, see “Business – Collaboration Arrangement – Collaboration Agreement With ImmuneOnco in Relation to the Development of IBC0966” in the Prospectus.
- *** We have completed Phase I clinical trials of relevant products as monotherapy, and plan to leverage data collected in the respective trials and directly seek IND approvals from competent regulatory authorities to conduct Phase II clinical trials of relevant products as combination therapy.

For further details of the product candidates of the Group, please refer to the Prospectus.

Warning: There is no assurance that we will ultimately be able to develop and market our Core Products or any of our pipeline products successfully.

Our Platforms

Our commitment to innovation is evident and supported by our proprietary technology platforms, which include (i) AIC™ Platform, a scalable platform mainly concentrated on antibody-cytokine fusion protein development, (ii) ADCC Enhanced Antibody Platform (“**AEA™ Platform**”), a FUT8 knock-out cell line constructed to enhance the cytotoxicity of antibodies, and (iii) Armed Innate Effector Multi-specific Platform (“**AIM™ Platform**”), a platform that focuses on the development of innate immunity stimulator-based bispecific/multi-specific antibodies. Each of them is designed for addressing technical difficulties and drug resistance faced in developing immunotherapies and achieving optimized treatment effects. Since their launch, we have developed IAP0971, IAE0972, IBB0979 and ISH0613 based on AIC™ Platform, IAH0968 based on AEA™ Platform, and IAN0982 based on AIM™ Platform.

AIC™ Platform

Our AIC™ Platform is prominently positioned in the field of immunocytokine development from multiple aspects, including cytokine selection and optimization, antibody selection and engineering, structural design and engineering, and production through customized cell line. It is a comprehensive research engine that includes not only a pool of intact immunoglobulin G antibodies and cytokines, but also functional antibody fragments and other types of immune system modulators. It is able to generate products ranging from immunocytokines to other bifunctional fusion proteins. Our clinical stage drug candidates IAP0971, IAE0972 and IBB0979, and preclinical stage drug candidate ISH0613 were developed based on the AIC™ Platform.

Management Discussion and Analysis

Core competencies of our AIC™ Platform include mechanism of action (“**MoA**”)-based antibody-cytokine selection, biology-oriented structural design and protein engineering, and production through customized cell lines.

- MoA-based antibody-cytokine selection is the cornerstone to achieve desired synergistic effects between antibody and cytokine. For example, selection of anti-PD-1 antibody and IL-15 cytokine for developing IAP0971 is grounded on their shared action site on the same T/NK cells, leading to great cis-synergy. The combination of anti-EGFR antibody and IL-10 is selected based on the potential engager effects it can produce. Specifically, IAE0972 can engage CD8+ T cells through IL-10 while simultaneously targeting tumor cells through the EGFR antibody moiety.
- Structural design and protein engineering module enable us to structurally design and modify our products to achieve improved safety and efficacy profile while reducing manufacturing cost and enhancing product quality manageability. Structural modifications that we are capable to perform through our AIC™ Platform include antibody and cytokine engineering, deglycosylation, linker/spacer design and optimization, and tertiary structure alteration.
- Production through customized cell lines is another important function performed by our AIC™ Platform. The cell lines we constructed for producing immunocytokines and other bifunctional fusion proteins are obtained after undergoing multiple rounds of metabolic and growth optimization and are of high expression capacity and excellent purification yield. Coupled with unique cytokine-specific codon optimization, stably expressed vehicles with optimized expression cassettes and our high-throughput screening system, it is able to reach an expression level of 4g/L and one-step affinity chromatography purity of 86%.

AEA™ Platform

Our AEA™ Platform is a biologically engineered Chinese hamster ovary (“**CHO**”) cell line with the FUT8 knocked-out to generate antibodies with enhanced ADCC and improved antitumor activities. Through this bioengineering modification, the CHO cell line will not be able to catalyze the transfer of fucose residue from its donor to its target, and thus is not able to produce any antibody that carries fucose. Because absence of core fucose on the Fc region has been shown to increase the Fc region’s binding affinity (up to 100 times) to its receptor FcγRIIIa present on immune effector cells, fucose-negative antibodies are expected to have enhanced ADCC activities through better activating immune effector cells.

Comparing to other platforms that aim to achieve enhanced ADCC by removing fucose from antibodies, AEA™ Platform is expected to produce antibodies with 0% of fucose, which stably and thoroughly enhances the ADCC of antibodies and simplifies quality control of the products.

Management Discussion and Analysis

AIM™ Platform

Our AIM™ Platform focuses on designing multi-functional biological products by engaging the innate immune system for cancer immunotherapy. It selects tumor associated antigen antibodies for cancer targeting, receptors agonist antibodies for innate effector activation, and cytokines and other TME factors for immune modulation to design multi-specific antibody fusion proteins, and evaluates them in terms of expression, target binding, in vitro and in vivo biological activities, as well as druggability. Currently, we have developed several categories of our proprietary AIM™ Platform that allow us to explore the combination of innate immunity stimulators with different types and numbers of targets, which provide us with abundant flexibility and diversity of various types of TME modulations for different clinical indications.

R&D

We consistently devote resources to R&D to pave for long-term growth. We believe the diversification and expansion of our product pipeline through both in-house R&D and through external collaboration are critical to our long-term competitiveness and success. Our fully-integrated biological therapeutic platform encompasses all the key biologic drug development functionalities, enabling us to identify and address potential clinical and manufacturing needs early in the development process, so we can direct our efforts towards biologics with best potential. Our platform spans from the early phase of identifying demand, developing core technologies, managing clinical trials, to the manufacturing of products. We believe that our integrated capabilities give us the agility to formulate our innovation, registration, commercialization and product optimization strategies that can navigate us through changing market needs, enable us to improve pipeline viability and expedite product development cycle at lower cost.

Collaboration Arrangement

In October 2019, we entered into a collaboration agreement (the “**IBC0966 Agreement**”) with ImmuneOnco with respect to the technology transfer, development, manufacture and commercialization of IBC0966. Pursuant to the IBC0966 Agreement, ImmuneOnco transferred to us (i) all of its rights and interests, including but not limited to development, production, regulatory filings and commercialization, in relation to IBC0966 in Mainland China, Hong Kong, Macau and Taiwan (the “**Territory**”); (ii) all related patents, if applicable, registered in the Territory; and (iii) all technical data and analytical methods relating to the development of IBC0966. Accordingly, ImmuneOnco has transferred to us its invention patent in Mainland China in relation to IBC0966 (patent number: CN111278865B), which covered all the key characteristics of IBC0966, and we have completed the administrative registration of the transfer. The application of this patent was filed on October 24, 2018 and the patent will expire on October 24, 2038.

Management Discussion and Analysis

Manufacturing

We have established our own global GMP-compliant manufacturing facilities, which meet both clinical and commercial production demands to quantity, quality and dosage form of our product candidates. We currently have four active drug substance production lines up to a total capacity of 1,600L, including three 200L and one 1,000L disposable bioreactors. We have successfully completed over 30 production batches of immunocytokines, mAbs, bsAbs and fusion proteins, which fulfilled the needs for performing preclinical studies, pilot production of antibody drugs and conducting early phase clinical trials. We have completed the installation of a production line for 5,000L bioreactor capacity, and completed the qualification in November 2023. When putting into operation, it will enable us to manufacture our drug candidates for Phase III clinical trials and commercialization in-house. Our drug product facility includes one commercial-scale liquid injection filling production line and one commercial scale lyophilized powder production line, which enables us to prepare biological products into various dosage forms according to different needs.

Future and Outlook

We plan to implement the following strategies to achieve our goals and visions:

- Focus on the development of immunocytokines to enhance position in this drug development field;
- Continue advancing selected pipeline products with great clinical value and commercial potential;
- Expand our GMP-compliant manufacturing facility to enhance our production capabilities and start to assemble our commercial team;
- Actively seek international collaboration opportunities to maximize value of our assets and increase brand awareness on a global scale; and
- Continue to focus on selecting and retaining top talents to fuel our innovation.

Management Discussion and Analysis

Financial Review

The following discussion is based on and should be read in conjunction with the financial information and accompanying notes included elsewhere in this report.

Other Income

During the Reporting Period and the six months ended June 30, 2024, other income consisted of (i) government grants by the PRC local government authorities mainly to support our R&D activities; and (ii) interest income from financial institutions. The following table sets forth a breakdown of our other income for the Reporting Period and the six months ended June 30, 2024:

	Six months ended June 30,	
	2025	2024
	RMB'000	RMB'000
Government grants	17	38
Interest income from financial institutions	5,018	1,744
Total	5,035	1,782

Other income of the Group increased by 178% from approximately RMB1.8 million for the six months ended June 30, 2024 to approximately RMB5.0 million for the Reporting Period, which was primarily due to the increase in interest income from financial institutions for the Reporting Period.

Other Gains and Losses, Net

During the Reporting Period, our other losses, net amounted to approximately RMB1.0 million, while our other gains, net for the six months ended June 30, 2024 was approximately RMB38.7 million. Our other gains and losses, net mainly consisted of (i) realized gains on other financial assets measured at FVTPL, mainly representing gains on the wealth management products we purchased; (ii) gains from fair value change of financial liabilities at FVTPL, mainly representing fair value gains of the preferred shares issued to the pre-IPO investors of our Company's global offering; and (iii) net foreign exchanges losses or gains. We recorded other losses, net for the six months ended June 30, 2025, primarily resulted from the net foreign exchange losses we recorded for the six months ended June 30, 2025.

R&D Expenses

During the Reporting Period and the six months ended June 30, 2024, our R&D expenses consisted of (i) contract research expenses in relation to the engagement of contract service providers; (ii) staff costs incurred by our R&D personnel; (iii) depreciation and amortization expenses in relation to our R&D machinery and equipment; (iv) material consumed in the course of our R&D activities; (v) application fees for our patents and IND applications; (vi) Share-based compensation; and (vii) other R&D expenses, mainly comprising traveling and transportation expenses of our R&D personnel, utilities incurred for our R&D activities and other miscellaneous expenses.

Management Discussion and Analysis

The following table sets forth a breakdown of our R&D expenses for the periods indicated.

	Six months ended June 30,	
	2025	2024
	RMB'000	RMB'000
Contract research expenses	5,384	4,559
Staff costs	8,176	7,614
Depreciation and amortization expenses	3,863	3,845
Materials consumed	2,053	1,447
Application fees	655	465
Share-based compensation	6,900	18,189
Others	1,736	1,589
Total	28,767	37,708

The R&D expenses decreased from approximately RMB37.7 million in the first half of 2024 to approximately RMB28.8 million during the Reporting Period, which was mainly due to the decrease in Share-based compensation of the Group in the first half of 2025 compared to that in the first half of 2024.

Administrative Expenses

During the Reporting Period and the six months ended June 30, 2024, our administrative expenses amounted to approximately RMB12.7 million and approximately RMB15.3 million, respectively, consisted of (i) general office expenses mainly comprising office product expenses, conference expenses and traveling and transportation expenses of administrative personnel; (ii) employee benefits expenses mainly relating to salaries, bonus and other welfare for our administrative employees; (iii) depreciation and amortization expenses for assets which were used for administrative purpose; (iv) professional service fees, which were primarily for related consulting, auditing and asset valuation in relation to corporate administration and restructuring; (v) Share-based compensation; and (vi) other administrative expenses mainly including tax and surcharges and other miscellaneous expenses. The decrease in administrative expenses for the Reporting Period as compared to that for the six months ended June 30, 2024 was primarily due to the decrease in administrative expenses and Share-based compensation related to the listing of the Company upon its completion.

Finance Costs

During the Reporting Period and the six months ended June 30, 2024, our finance costs amounted to approximately RMB0.8 million and approximately RMB0.6 million, respectively, which consisted of (i) interest expenses on bank borrowings; and (ii) interest expenses on our lease liabilities. The increase in finance costs for the Reporting Period as compared to that for the six months ended June 30, 2024 was primarily due to the increase in interest expenses on bank borrowings.

Management Discussion and Analysis

Listing Expenses

Listing expenses represent expenses incurred for our listing and global offering. During the Reporting Period, we did not record any listing expenses. For the six months ended June 30, 2024, we recorded listing expenses of approximately RMB23.0 million.

Income Tax Expenses

Our income tax expense for the Reporting Period was nil (six months ended June 30, 2024: nil).

Loss for the Period

As a result of the foregoing, our loss for the period increased from approximately RMB36.1 million for the six months ended June 30, 2024 to approximately RMB38.2 million for the Reporting Period.

Liquidity and Financial Resources

We have continued to maintain a healthy and sound financial position and have followed a set of funding and treasury policies to manage our capital resources and mitigate potential risks involved. As of June 30, 2025, the Group's total cash and cash equivalents amounted to approximately RMB171 million, representing an increase of approximately 116% as compared to approximately RMB79.0 million as of December 31, 2024. Such increase was primarily due to the maturity of time deposits.

As of June 30, 2025, the balance of the time deposits of the Group was approximately RMB238.4 million. As of December 31, 2024, the time deposits of the Group was approximately RMB219.5 million.

As of June 30, 2025, current assets of the Group amounted to approximately RMB488.1 million; and current liabilities of the Group amounted to approximately RMB74.1 million, including interest-bearing bank borrowings of approximately RMB61.3 million. Bank borrowings of our Group were denominated in RMB, and were secured and credit loans, payable within 12 months and carried an annual interest rate ranging from 3.22% to 3.44%.

Management Discussion and Analysis

Indebtedness

The following table sets forth the breakdown of our bank loans, lease liabilities and financial liabilities at FVTPL as of the dates indicated:

	As of June 30, 2025 RMB'000	As of December 31, 2024 RMB'000
Secured and unguaranteed		
Bank loans	41,300	9,500
Lease liabilities	–	22
Unsecured and unguaranteed		
Lease liabilities	6,944	6,874
Financial liabilities at FVTPL	–	–
Bank loans	20,000	24,800
Total	68,244	41,196

Save as discussed above, we did not have any other material mortgages, charges, debentures, loan capital, debt securities, loans, bank overdrafts or other similar indebtedness, finance lease or hire purchase commitments, liabilities under acceptance (other than normal trade bills), acceptance credits, which are either guaranteed, unguaranteed, secured or unsecured, or guarantees or other contingent liabilities as of June 30, 2025.

Gearing Ratio

As of June 30, 2025, the gearing ratio, calculated by dividing total liabilities by total assets and multiplied by 100%, increased to approximately 13.73%, as compared with approximately 8.7% as of December 31, 2024.

Significant Investments, Material Acquisitions and Disposal

The Group did not have any significant investments or material acquisitions or disposals of subsidiaries, associates and joint ventures for the six months ended June 30, 2025.

Capital Commitments

As of June 30, 2025, we had capital commitment of RMB19.1 million, primarily arose from the contracts we entered into with suppliers for the acquisition of equipment and the contract we entered into to acquire the land use right to support the construction of our production lines and the expansion of our business operations (as of December 31, 2024: RMB23.3 million).

Management Discussion and Analysis

Contingent Liabilities

Except for the under provision of social insurance and housing provident fund contributions, we did not have any material contingent liabilities as of December 31, 2024 and June 30, 2025. For the related risk, see “Risk Factors – Risks Relating to Our Operations – Any failure to comply with the PRC regulations regarding contribution of social insurance premium or housing provident funds may subject us to fines and other legal or administrative measures” in the Prospectus.

Pledge of Assets

The bank loans of RMB41,300,000 as of June 30, 2025 were secured, unguaranteed and carried fixed interest rate ranging from 3.22% to 3.44% (as of December 31, 2024: RMB9,500,000; 3.44%). Such bank loans were secured by bank deposits of USD6,365,000 (equivalent to approximately RMB45,564,489).

Foreign Exchange Exposure

Foreign currency risk refers to the risk of loss resulting from changes in foreign currency exchange rates. Fluctuations in exchange rates between RMB and other currencies in which our Group conducts business may affect our financial condition and results of operation. The Group mainly operates in the PRC and is exposed to foreign exchange risk arising from various currency exposures, primarily with respect to HK\$ and USD. The conversion of foreign currencies into RMB, including HK\$ and the USD, has been based on rates set by the People’s Bank of China. The Group primarily limits our exposure to foreign currency risk by closely monitoring the foreign exchange market. During the Reporting Period, the Group did not enter into any currency hedging transactions.

Future Plan for Material Investments or Capital Assets

As of June 30, 2025, save as disclosed in this report, the Group did not have any future plan for material investments or capital assets.

Material Acquisitions and Disposals of Subsidiaries, Associates and Affiliated Companies

The Group did not have any material acquisitions or disposals of subsidiaries, associates and joint ventures during the Reporting Period.

Employee and Remuneration

As of June 30, 2025, our Group had a total of 128 employees. The total remuneration cost of our Group for the Reporting Period was approximately RMB18.1 million, as compared to approximately RMB28.5 million for the six months ended June 30, 2024. We have designed an evaluation system to assess the performance of its employees periodically. Such system forms the basis of our determinations of whether an employee should receive a salary raise, bonus, or promotion. We believe the salaries and bonuses that the employees receive are competitive with market rates.

We place strong emphasis on providing training to our employees in order to enhance their technical and product knowledge. We design and offer different training programmes for our employees in various positions.

Management Discussion and Analysis

We make contributions to the social insurance and housing provident fund for all of our employees in the PRC. We have adopted the RSU Scheme to recognize and motivate the contributions by the relevant participants and give incentives thereto in order to retain them, as well as to attract suitable personnel for further development of our Group. Please refer to “D. RSU Scheme” in Appendix IV to the Prospectus for a summary of the principal terms of the RSU Scheme.

Use of Proceeds

The Group received net proceeds (after deduction of underwriting commissions and related costs and expenses) from our Company’s global offering of approximately HK\$391.6 million. The net proceeds from our Company’s global offering have been and will be used in accordance with the purposes as set out in the Prospectus. The following table sets forth the use of the net proceeds from our Company’s global offering as of June 30, 2025:

Proposed use of proceeds	Allocation of net proceeds from the Global Offering (HK\$ million)	Percentage of total net proceeds (%)	Utilized amount (as of June 30, 2025) (HK\$ million)	Unutilized amount (as of June 30, 2025) (HK\$ million)	Expected timeline of full utilization of the remaining net proceeds as of June 30, 2025
For ongoing and planned clinical trials of IAH0968 in China	110.4	28.2	51.8	58.6	By the end of 2028
For ongoing and planned clinical trials of IAP0971 in China	140.1	35.8	15.8	124.3	By the end of 2028
For ongoing and planned clinical trials of IAE0972 in China	141.1	36.0	8.0	133.1	By the end of 2028
Total	391.6	100	75.6	316.0	

As of the date of this report, there is no material change in the use of the proceeds from the Company’s global offering as stated in the Prospectus. Based on the R&D progress of our Core Products, the Company expects that the net proceeds from the Company’s global offering will be used up by the end of 2028.

Events after the Reporting Period

Deloitte Touche Tohmatsu resigned as the auditor of the Company with effect from September 9, 2025. ZHONGHUI ANDA CPA Limited was appointed as the new auditor of the Company with effect from September 16, 2025. For details, please refer to the announcements of the Company dated September 15, 2025 and September 16, 2025.

Save as disclosed above, there has been no important event subsequent to the Reporting Period and up to the date of this report, which would affect the Group’s business operations in material aspects.

Other Information

Updates on Information of Directors

There was no change in the Board and the information of Directors between the date of the Company's 2024 annual report and the date of this report, which is required to be disclosed pursuant to Rule 13.51B(1) of the Listing Rules.

Directors' and Chief Executive's Interests and Short Positions in Shares, Underlying Shares and Debentures of the Company and its Associated Corporations

As at June 30, 2025, the interests or short positions of the Directors and chief executive of the Company in the shares, underlying shares and debentures of the Company or any of its associated corporations (within the meaning of Part XV of the SFO), which (a) were required to be notified to the Company and the Stock Exchange pursuant to Divisions 7 and 8 of Part XV of the SFO (including interests and short positions which he/she was taken or deemed to have under such provisions of the SFO); or (b) were required, pursuant to section 352 of the SFO, to be recorded in the register referred to therein; or (c) were required to be notified to the Company and the Stock Exchange pursuant to the Model Code, were as follows:

Name of Director or chief executive	Capacity/Nature of interest	Number of Shares as at June 30, 2025 ⁽¹⁾	Approximate percentage of the Company's issued share capital ⁽²⁾
Mr. ZHANG Feng (張峰) ⁽³⁾	Interests in controlled corporations	100,000,000	63.83%
Ms. JIANG Xiaoling (姜曉玲) ⁽⁴⁾	Beneficial owner	500,000	0.32%

Notes:

(1) All interest stated are long positions.

(2) The calculation is based on the total number of 156,666,800 Shares in issue as at June 30, 2025.

(3) Sunho Wisdom is owned as to 99.9% by Sunho Fortune (as a nominee which is wholly owned by a trust established by Mr. ZHANG Feng as the settlor and beneficiary) and 0.1% by Innovalue Investments (a wholly-owned subsidiary of Mr. ZHANG Feng), respectively. Further, Mr. ZHANG Feng is entitled to exercise approximately 73.19% voting rights in No5XJR through Innovalue Investments. Sunho Stellar is wholly owned by an independent professional trustee who shall exercise all voting rights attached to the Shares held by Sunho Stellar in accordance with the instructions of Mr. ZHANG Feng. As such, under the SFO, Mr. ZHANG Feng is deemed to be interested in the Shares held by Sunho Wisdom, No5XJR and Sunho Stellar.

(4) These Shares represent the entitlement of Ms. JIANG Xiaoling (姜曉玲) to receive up to 500,000 Shares pursuant to the RSUs granted to her under the RSU Scheme, subject to the terms and conditions of these RSUs.

Other Information

Save as disclosed above and to the best knowledge of the Directors, as of June 30, 2025, none of the Directors or the chief executive of the Company has any interests and/or short positions in the shares, underlying shares or debentures of the Company or its associated corporations (within the meaning of Part XV of the SFO) which were required to be notified to the Company and the Stock Exchange pursuant to Divisions 7 and 8 of Part XV of the SFO (including interests and short positions which they are taken or deemed to have under such provisions of the SFO), or which were required, pursuant to section 352 of the SFO, to be entered in the register referred to therein, or which were required, pursuant to the Model Code, to be notified to the Company and the Stock Exchange.

Substantial Shareholders' Interests

So far as our Directors are aware, as at June 30, 2025, the following persons have interests or short positions in Shares or underlying Shares of our Company which will be required to be disclosed to our Company and the Stock Exchange pursuant to the provisions of Divisions 2 and 3 of Part XV of the SFO and recorded in the register required to be maintained by the Company under Section 336 of the SFO:

Name of Shareholder	Capacity/Nature of interest	Number of Shares as at June 30, 2025 ⁽¹⁾	Approximate percentage of the Company's issued share capital ⁽²⁾
Sunho Wisdom	Beneficial owner	88,000,000	56.17%
Sunho Fortune ⁽³⁾	Interests in controlled corporations	88,000,000	56.17%
Trident Trust Company (HK) Limited ⁽³⁾	Trustee	94,000,000	60.00%
Mr. ZHANG Feng (張峰) ⁽³⁾	Interests in controlled corporations	100,000,000	63.83%
Huzhou Efung Ansheng Venture Capital Partnership (Limited Partnership) (湖州市倚鋒安盛創業投資合夥企業 (有限合夥)) ("Efung Ansheng")	Beneficial owner	11,666,660	7.45%
Shenzhen Efung Investment Management Enterprise (Limited Partnership) (深圳市倚鋒投資管理企業(有限合夥)) ⁽⁴⁾	Interests in controlled corporations	11,666,660	7.45%
Shenzhen Efung Venture Capital Investment Co., Ltd. (深圳市倚鋒創業投資有限公司) ⁽⁴⁾	Interests in controlled corporations	11,666,660	7.45%
Shenzhen Efung Holdings Group Co., Ltd. (深圳市倚鋒控股集團有限公司) ("Efung Holdings") ⁽⁴⁾	Interests in controlled corporations	17,500,000	11.17%
Mr. ZHU Jinqiao (朱晉橋) ⁽⁴⁾	Interests in controlled corporations	17,500,000	11.17%
Guocheng (Zhejiang) Industrial Development Co., Ltd. (國成(浙江)實業發展有限公司) ⁽⁴⁾	Interests in controlled corporations	17,500,000	11.17%

Other Information

Substantial Shareholders' Interests (Continued)

Notes:

- (1) All interest stated are long positions.
- (2) The calculation is based on the total number of 156,666,800 Shares in issue as at June 30, 2025.
- (3) Sunho Wisdom is owned as to 99.9% by Sunho Fortune (as a nominee which is wholly owned by a trust established by Mr. ZHANG Feng as the settlor and beneficiary) and 0.1% by Innovalue Investments (a wholly-owned subsidiary of Mr. ZHANG Feng), respectively. As such, under the SFO, Sunho Fortune is deemed to be interested in the Shares held by Sunho Wisdom. Further, Mr. ZHANG Feng is entitled to exercise approximately 73.19% voting rights in No5XJR through Innovalue Investments. Sunho Stellar is wholly owned by an independent professional trustee who shall exercise all voting rights attached to the Shares held by Sunho Stellar in accordance with the instructions of Mr. ZHANG Feng. As such, under the SFO, Mr. ZHANG Feng is deemed to be interested in the Shares held by Sunho Wisdom, No5XJR and Sunho Stellar.
- (4) Efung Ansheng is a limited partnership established in the PRC and is managed by its general partner, Shenzhen Efung Investment Management Enterprise (Limited Partnership) (深圳市倚鋒投資管理企業(有限合夥)), whose general partner is Shenzhen Efung Venture Capital Investment Co., Ltd. (深圳市倚鋒創業投資有限公司) which is in turn held as to approximately 60% by Efung Holdings and approximately 40% by Mr. ZHU Jinqiao (朱晉橋). As such, each of Shenzhen Efung Investment Management Enterprise (Limited Partnership) and Shenzhen Efung Venture Capital Investment Co., Ltd. is deemed to be interested in the Shares held by Efung Ansheng. Huzhou Efung Anhe Venture Capital Partnership (Limited Partnership) (湖州市倚鋒安禾創業投資合夥企業(有限合夥)) ("Efung Anhe") is a limited partnership established in the PRC and is managed by its general partner, Hainan Efung Junma Private Equity Fund Management Co., Ltd. (海南倚鋒駿馬私募基金管理有限公司), which is held as to approximately 70% by Efung Holdings. Efung Holdings is held as to approximately 54% by Mr. ZHU Jinqiao (朱晉橋). Besides, Guocheng (Zhejiang) Industrial Development Co., Ltd. (國成(浙江)實業發展有限公司) holds approximately 99.99% partnership interest in Efung Ansheng as its limited partner and approximately 49.99% partnership interest in Efung Anhe as its limited partner. As such, each of Efung Holdings, Mr. ZHU Jinqiao and Guocheng (Zhejiang) Industrial Development Co., Ltd. is deemed to be interested in the Shares held by Efung Ansheng and Efung Anhe.

Save as disclosed above and to the best knowledge of the Directors, as of June 30, 2025, no person (other than the Directors or chief executives of the Company) had an interest or a short position in the shares or underlying shares of the Company as recorded in the register of interest required to be kept by the Company under section 336 of the SFO.

RSU SCHEME

On August 2, 2023, the Company adopted the RSU Scheme. The purpose of the RSU Scheme is to recognize and motivate the contributions by the Participants (as defined below) and give incentives thereto in order to retain them, as well as to attract suitable personnel for further development of our Group. An award of RSU(s) under the RSU Scheme (the "**Award**") gives a Participant (as defined below) a conditional right upon vesting of the Award to obtain either Share(s) or an equivalent value in cash with reference to the value of the Share(s) on or about the date of vesting, as determined by our Board at its absolute discretion, less any tax, fees, levies, stamp duty and other applicable charges. An Award may include, if so specified by our Board in its entire discretion, cash and non-cash income, dividends or distributions and/or the sale proceeds of non-cash and non-scrip distributions in respect of those Shares underlying the RSU(s) from the date that the Award is granted to the date that it vests. Unless otherwise duly approved by our Shareholders, the total number of Shares underlying the RSU Scheme shall not exceed 6,000,000 Shares, subject to any adjustment pursuant to any reorganization of capital structure from time to time. Participants of the RSU Scheme include employees or officers of our Group, including executive, non-executive and independent non-executive directors of our Group and any prospective employees who receive a grant as an inducement to join our Group (the "**Participant(s)**").

Other Information

Details of movements of the RSUs granted under the RSU Scheme during the six months ended June 30, 2025 are set out below:

Participants	Position	Date of Grant	Vesting period	Exercise price	Number of Shares underlying the RSUs granted	Number of RSUs outstanding as at January 1, 2025	Number of RSUs granted during the Reporting Period	Number of RSUs vested during the Reporting Period	Number of RSUs cancelled during the Reporting Period	Number of RSUs lapsed during the Reporting Period	Weighted average closing price immediately before the dates of vesting during the Reporting Period (HK\$)	Number of outstanding RSUs as at June 30, 2025
Ms. JIANG Xiaoling (姜曉玲)	Executive Director and vice president	May 6, 2023 ⁽¹⁾	Note ⁽²⁾	nil	500,000	500,000	-	(100,000)	-	-	5.28	400,000
Ms. XU Chunqin (徐春芹)	Chief financial officer and joint company secretary	May 6, 2023 ⁽¹⁾	Note ⁽²⁾	nil	500,000	500,000	-	(100,000)	-	-	5.28	400,000
Mr. JIANG Dongcheng (姜東成)	Vice president	May 6, 2023 ⁽¹⁾	Note ⁽²⁾	nil	5,000,000	5,000,000	-	(1,000,000)	-	-	5.28	4,000,000
Total						6,000,000	-	(1,200,000)	-	-	5.28	4,800,000

Notes:

- (1) The above grantees were granted share incentives on May 6, 2023, the terms of which (including the performance targets) were amended upon the formal adoption of the RSU Scheme on August 2, 2023.
- (2) 20% of the RSUs granted shall vest on each of the first, second, third, fourth and fifth anniversary of the Listing Date, provided that certain conditions (including certain performance targets of our Group and certain performance ratings of the grantees) are met.

As at June 30, 2025, the number of Awards available for grant under the RSU Scheme was nil (as at January 1, 2025: nil).

The number of Shares that may be issued in respect of Awards granted under the RSU scheme during the Reporting Period divided by the weighted average number of Shares in issue (excluding treasury shares) for the Reporting Period was nil.

Continuing Disclosure Obligation Pursuant to the Listing Rules

Save as disclosed in this report, the Company does not have any other disclosure obligations under Rules 13.20, 13.21 and 13.22 of the Listing Rules.

Compliance with the Corporate Governance Code

The Company has adopted the principles and code provisions in the Corporate Governance Code ("CG Code") set out in Appendix C1 to the Listing Rules and has complied with all applicable code provisions of the CG Code during the Reporting Period.

Model Code for Securities Transactions

The Company has adopted the Model Code as its own code of conduct regarding securities transactions by the Directors. Having made specific enquiries with all Directors, each of them has confirmed that he/she has complied with the Model Code during the Reporting Period. No incident of non-compliance of the Model Code by the employees who are likely to be in possession of inside information of the Company was noted by the Company.

Other Information

Purchase, Sale or Redemption of Listed Securities

Neither the Company nor any of its subsidiaries purchased, sold or redeemed any of the Company's listed securities during the Reporting Period (including sale of treasury shares, if any). As of the date of this report, the Company did not hold any treasury shares (as defined in the Listing Rules).

Audit Committee

The Company has established the Audit Committee with written terms of reference in compliance with the CG Code. As at the date of this report, the Audit Committee consists of three independent non-executive Directors, namely Mr. CHAN Heung Wing Anthony, Ms. FENG Lan and Mr. SHI Luwen. Mr. CHAN Heung Wing Anthony is the chairman of the Audit Committee, who possesses suitable professional qualifications.

Review of Interim Results

The Audit Committee has reviewed this interim report and the unaudited interim financial information for the Reporting Period (with no disagreement), together with the management of the Company. The Audit Committee has also reviewed the accounting principles and practices adopted by the Group and discussed, risk management, internal control and financial reporting matters of the Group for the Reporting Period. There is no disagreement between the Board and the Audit Committee regarding the accounting treatment adopted by the Company.

Interim Dividend

The Board does not recommend the payment of an interim dividend for the Reporting Period (for the six months ended June 30, 2024: nil).

Condensed Consolidated Statements of Profit or Loss and Other Comprehensive Income

For the six months ended June 30, 2025

	Notes	Six months ended June 30,	
		2025	2024
		RMB'000	RMB'000
		(unaudited)	(unaudited)
Other income	5	5,035	1,782
Other gains and losses, net	6	(963)	38,720
Research and development expenses		(28,767)	(37,708)
Administrative expenses		(12,678)	(15,270)
Listing expenses		—	(23,035)
Finance costs	7	(826)	(566)
Loss before tax		(38,199)	(36,077)
Income tax expense	8	—	—
Loss and total comprehensive expense for the period	9	(38,199)	(36,077)
Loss per share			
— Basic and diluted (RMB per share)	11	(0.25)	(0.34)

Condensed Consolidated Statements of Financial Position

As at June 30, 2025

	Notes	June 30, 2025 RMB'000 (unaudited)	December 31, 2024 RMB'000 (audited)
Non-current assets			
Property and equipment	12	31,972	34,812
Right-of-use assets	12	15,759	16,992
Intangible asset		10,000	10,000
Equity instrument at FVTOCI		910	910
Prepayments for acquisition of equipment		8,675	2,523
Refundable fulfillment deposits	13	2,500	2,500
		69,816	67,737
Current assets			
Inventories		1,101	974
Deposits, prepayments and other receivables	13	31,854	24,231
Financial assets at fair value through profit or loss ("FVTPL")		–	158,825
Time deposits		238,381	219,468
Restricted bank deposits		45,564	10,509
Cash and cash equivalents		171,166	78,991
		488,066	492,998
Current liabilities			
Trade and other payables	14	8,372	7,601
Bank loans	15	61,300	34,300
Lease liabilities		4,403	2,245
		74,075	44,146
Net current assets		413,991	448,852
Total assets less current liabilities		483,807	516,589

Condensed Consolidated Statements of Financial Position

As at June 30, 2025

	Note	June 30, 2025 RMB'000 (unaudited)	December 31, 2024 RMB'000 (audited)
Non-current liabilities			
Lease liabilities		2,541	4,651
		2,541	4,651
Net assets		481,266	511,938
Capital and reserves			
Share capital	16	524	524
Treasury stock		(15)	(19)
Reserves		480,757	511,433
Total equity		481,266	511,938

Condensed Consolidated Statements of Changes in Equity

For the six months ended June 30, 2025

	Attributable to owners of the Company						Total RMB'000
	Share capital RMB'000	Treasury stock RMB'000	Capital reserve RMB'000	Share premium RMB'000	Share-based payment reserve RMB'000	Accumulated losses RMB'000	
As at January 1, 2024 (audited)	322	(19)	223,704	–	–	(328,116)	(104,109)
Loss and total comprehensive expense for the period	–	–	–	–	–	(36,077)	(36,077)
Recognition of equity-settled share-based payments expenses (note 17)	–	–	–	–	19,843	–	19,843
Issue of shares upon initial public offering ("IPO") (note 16)	122	–	–	419,655	–	–	419,777
Share issue costs for IPO	–	–	–	(28,154)	–	–	(28,154)
Automatic conversion of Series A Preferred Shares upon IPO (note 16)	80	–	–	276,663	–	–	276,743
As at June 30, 2024 (unaudited)	524	(19)	223,704	668,164	19,843	(364,193)	548,023
As at January 1, 2025 (audited)	524	(19)	223,704	667,462	28,348	(408,081)	511,938
Loss and total comprehensive expense for the period	–	–	–	–	–	(38,199)	(38,199)
Recognition of equity-settled share-based payments expenses (note 17)	–	–	–	–	7,527	–	7,527
Shares vested in connection with RSU Plan (note 17)	–	4	–	11,726	(11,730)	–	–
As at June 30, 2025 (unaudited)	524	(15)	223,704	679,188	24,145	(446,280)	481,266

Condensed Consolidated Statements of Cash Flows

For the six months ended June 30, 2025

	Six months ended June 30, 2025 RMB'000 (unaudited)	2024 RMB'000 (unaudited)
NET CASH USED IN OPERATING ACTIVITIES	(36,115)	(120,198)
INVESTING ACTIVITIES		
Interest received from banks	5,310	720
Acquisition of property and equipment	(6,803)	(5,129)
Disposal of property and equipment	–	721
Acquisition of land use right	–	(9,600)
Purchase of other financial assets	–	(2,132)
Purchase of financial assets at FVTPL	–	(910)
Redemption of FVTPL	158,825	51,711
Interest received from other financial assets	–	1,022
Redemption of time deposits with maturity of more than three months	10,509	35,414
Interest received from time deposits with maturity of more than three months	–	1,024
Placement of restricted bank deposits	(45,564)	–
Placement from time deposits with maturity of more than three months	(18,913)	–
NET CASH FROM INVESTING ACTIVITIES	103,364	72,841
FINANCING ACTIVITIES		
New loan raised	41,800	22,980
Repayment of bank loans	(14,800)	–
Interest paid on bank loans	(778)	–
Proceeds from issuance of ordinary shares	–	419,777
Issue costs paid	–	(24,214)
Payment of lease liabilities	–	(2,378)
NET CASH FROM FINANCING ACTIVITIES	26,222	416,165
NET INCREASE IN CASH AND CASH EQUIVALENTS	93,471	368,808
Effect of foreign exchange rate changes	(1,296)	–
CASH AND CASH EQUIVALENTS AT THE BEGINNING OF THE PERIOD	78,991	125,074
TOTAL CASH AND CASH EQUIVALENTS AT THE END OF THE PERIOD, represented by cash and cash equivalents	171,166	493,882

Notes to the Condensed Consolidated Financial Statements

For the six months ended June 30, 2025

1. General Information

Sunho Biologics, Inc. (the “**Company**”) was incorporated in the Cayman Islands as an exempted company registered under the Company Laws of the Cayman Islands on May 14, 2021. The shares of the Company have been listed on the Main Board of The Stock Exchange of Hong Kong Limited (“**Stock Exchange**”) with effect from May 24, 2024. The Company’s ultimate controlling shareholder is Mr. Zhang Feng who achieves ultimate control through his direct or indirect interests held in the Company. The address of the registered office is PO Box 309, Ugland House, Grand Cayman, KY1-1104, Cayman Islands, and the principal place of business of the Company is 31/F, Tower Two, Times Square, 1 Matheson Street, Causeway Bay, Hong Kong.

The Company is an investment holding company. The Company and its subsidiaries (collectively referred to as the “**Group**”) are mainly committed to the develop regulate immune microenvironment by directly modulating both the innate and adaptive immune systems.

The condensed consolidated financial statements are presented in Renminbi (“**RMB**”), which is the functional currency of the Company and its subsidiaries.

2. Basis of Preparation

The condensed consolidated financial statements have been prepared in accordance with International Accounting Standard 34 “Interim Financial Reporting” issued by the International Accounting Standards Board (the “**IASB**”) as well as the applicable disclosure requirements of the Rule Governing the Listing of Securities on the Stock Exchange.

3. Principal Accounting Policies

The condensed consolidated financial statements have been prepared on the historical cost basis except for certain financial instruments which are measured at fair values.

Other than additional/change in accounting policies resulting from application of amendments to IFRS Accounting Standards, the accounting policies and methods of computation used in the condensed consolidated financial statements for the six months ended June 30, 2025 are the same as those followed in the preparation of the Group’s annual consolidated financial statements for the year ended December 31, 2024.

Application of amendments to IFRS Accounting Standards

The IASB has issued a number of amendments to IFRS Accounting Standards that are first effective for the current accounting period of the Group. None of these developments has had a material effect on how the Group’s results and financial position for the current or prior periods have been prepared or presented in the interim financial report. The Group has not applied any new IFRS Accounting Standards that are not yet effective for the current accounting period.

Notes to the Condensed Consolidated Financial Statements

For the six months ended June 30, 2025

4. Segment Information

For the purposes of resources allocation and performance assessment, the Group's chief executive officer, being the chief operating decision maker ("CODM"), reviews the overall results and financial position of the Group as a whole when making decisions about allocating resources and assessing performance of the Group and hence, the Group has only one operating and reportable segment and no further analysis of this single segment is presented.

The Group did not record any revenue for the six months ended June 30, 2025 (six months ended June 30, 2024: nil). As at June 30, 2025, the Group's all non-current assets are located in the People's Republic of China ("PRC") and accordingly, no analysis of geographical information is presented.

5. Other Income

	Six months ended June 30,	
	2025	2024
	RMB'000	RMB'000
	(unaudited)	(unaudited)
Government grants (note)	17	38
Interest income from financial institutions	5,018	1,744
	5,035	1,782

Note: The amount represents subsidies granted by the PRC local government authorities as incentives for the Group's research and development activities. The government grants including unconditional and conditional and had been approved by the PRC local government authorities. The unconditional government grants are recognized when payments were received. The conditional government grants are recognized when condition met and the corresponding grants are received.

Notes to the Condensed Consolidated Financial Statements

For the six months ended June 30, 2025

6. Other Gains and Losses, Net

	Six months ended June 30,	
	2025	2024
	RMB'000	RMB'000
	(unaudited)	(unaudited)
Realized gain on other financial assets	–	1,022
Gain from fair value change of financial liabilities at FVTPL	–	34,782
Net foreign exchange (losses)/gains	(1,004)	2,880
Others	41	36
	(963)	38,720

7. Finance Costs

	Six months ended June 30,	
	2025	2024
	RMB'000	RMB'000
	(unaudited)	(unaudited)
Interest expenses on bank borrowings	778	447
Interest expenses on lease liabilities	48	119
	826	566

8. Income Tax Expense

	Six months ended June 30,	
	2025	2024
	RMB'000	RMB'000
	(unaudited)	(unaudited)
Current PRC enterprise income tax	–	–

The Company was incorporated in the Cayman Islands and Sunho bio Investments Limited was incorporated in the BVI that are tax exempted.

No Hong Kong profits tax was provided as there was no assessable profit that was subjected to Hong Kong Profits Tax during both periods.

Under the Law of the PRC on Enterprise Income Tax (the “EIT Law”) and Implementation Regulation of the EIT Law, the tax rate of the PRC subsidiaries is 25% for the six months ended June 30, 2025.

Notes to the Condensed Consolidated Financial Statements

For the six months ended June 30, 2025

8. Income Tax Expense (Continued)

Pursuant to Caishui 2023 circular No. 7, 盛禾(中國)生物製藥有限公司 Sunho (China) Biopharmaceutical Co., Ltd.* ("**Sunho (China) Biopharmaceutical**") enjoyed super deduction of 200% on qualified research and development expenditures during both periods.

As at June 30, 2025, the Group has no deductible temporary differences (December 31, 2024: nil). No deferred tax asset has been recognized in relation to such deductible temporary difference as it is not probable that taxable profit will be available against which the deductible temporary differences can be utilized.

* English name for identification purpose only.

9. Loss for the Period

	Six months ended June 30,	
	2025	2024
	RMB'000	RMB'000
	(unaudited)	(unaudited)
Loss for the period has been arrived at after charging:		
Depreciation of property and equipment	3,491	3,363
Depreciation of right-of-use assets	1,233	1,132
	4,724	4,495
Listing expenses	—	23,035
Directors' emoluments	1,963	2,682
Other staff costs:		
— salaries and other benefits	5,594	6,831
— retirement benefit scheme contribution	751	761
— share-based payments	6,900	18,189
	15,208	28,463

10. Dividends

No dividends were paid, declared or proposed during the interim period. The directors of the Company have determined that no dividend will be paid in respect of the interim period (six months ended June 30, 2024: nil).

Notes to the Condensed Consolidated Financial Statements

For the six months ended June 30, 2025

11. Loss Per Share

The calculation of the basic and diluted loss per share attributable to the owners of the Company is based on the following data:

	Six months ended June 30,	
	2025 (unaudited)	2024 (unaudited)
Loss (RMB'000)		
Loss for the period attributable to the owners of the Company for the purpose of calculating basic and diluted loss per share	(38,199)	(36,077)
Number of shares ('000)		
Weighted average number of ordinary shares for the purpose of calculating basic and diluted loss per share	150,919	105,520

The basic loss per share is calculated based on the loss attributable to the owners of the Company and the weighted average number of ordinary shares excluded shares of treasury stock under the employee incentive schemes (note 17).

Diluted loss per share is calculated by adjusting the weighted average number of ordinary shares outstanding to assume conversion of all dilutive potential ordinary shares. The computation of diluted loss per share for the six months ended June 30, 2025 does not assume the vesting of share-based awards granted to employees (note 17) (six months ended June 30, 2024: does not assume the conversion of the Series A Preferred Shares before IPO (Details of the Series A Preferred Shares before IPO were disclosed in note 27 of the Group's consolidated financial statements for the year ended December 31, 2024) and the vesting of share-based awards granted to employees (note 17)) since their assumed conversion or vesting would result in a decrease in loss per share. Accordingly, diluted loss per share for the periods ended June 30, 2025 and 2024 are the same as basic loss per share respectively.

Notes to the Condensed Consolidated Financial Statements

For the six months ended June 30, 2025

12. Property and Equipment and Right-of-use Assets

During the six months ended June 30, 2025, the Group acquired additional property, plant and equipment with an aggregate cost of RMB651,000 (six months ended June 30, 2024: nil).

During the six months ended June 30, 2024, the Group disposed of construction in progress with an aggregate carrying amount of RMB721,000 for proceeds of RMB721,000.

During the six months ended June 30, 2024, the Group incurred RMB9,600,000 on acquisition of leasehold land of which the Group is in the process of obtaining the land use right certificate.

During the current interim period, no addition lease agreement was entered by the Group. During the period ended June 30, 2024, no addition lease agreement was entered by the Group.

13. Deposits, Prepayments And Other Receivables

	As at June 30, 2025 RMB'000 (unaudited)	As at December 31, 2024 RMB'000 (audited)
Value added tax recoverable	5,528	4,860
Prepayments for research and development costs	22,462	15,082
Refundable fulfillment deposits	2,500	2,500
Interest receivables	979	1,271
Refundable tendering deposits	960	960
Others	1,925	2,058
	34,354	26,731
Analyzed as:		
Non-current	2,500	2,500
Current	31,854	24,231
	34,354	26,731

Notes to the Condensed Consolidated Financial Statements

For the six months ended June 30, 2025

14. Trade and Other Payables

	As at June 30, 2025 RMB'000 (unaudited)	As at December 31, 2024 RMB'000 (audited)
Payables for research and development costs	2,287	1,022
Accrued research and development costs	1,397	2,236
Accrued staff costs and benefits	2,471	2,151
Other payables:		
Payable for equipment	613	326
Accrued professional service fee	535	1,572
Others	1,035	229
Other tax payables	34	65
	8,372	7,601
Analyzed as:		
Current	8,372	7,601

The average credit period on purchases of materials and services of the Group is 10–60 days.

The following is an aging analysis of payables for payables for research and development costs, presented based on the invoice dates at the end of each reporting period:

	As at June 30, 2025 RMB'000 (unaudited)	As at December 31, 2024 RMB'000 (audited)
0–90 days	–	2
Over 90 days	2,287	1,020
	2,287	1,022

Notes to the Condensed Consolidated Financial Statements

For the six months ended June 30, 2025

15. Bank Loans

During the current interim period, the Group raised new bank loans amounting to RMB41,800,000 (six months ended June 30, 2024: RMB22,980,000). The loans carry interest at fixed market rates of 3.22% to 3.44% (December 31, 2024: 3.35% to 3.80%) and unguaranteed and are repayable in twelve months.

The bank loans of RMB41,300,000 as at June 30, 2025 (December 31, 2024: RMB9,500,000) are secured, unguaranteed and carried fixed interest rate of 3.22%-3.44% (December 31, 2024: 3.44%). Such bank loans are secured by bank deposits of USD6,365,000 (equivalent to approximately RMB45,564,000) (December 31, 2024: USD1,460,000 (equivalent to approximately RMB10,509,000)

16. Share Capital

The Company was incorporated in the Cayman Islands on May 14, 2021, with authorized share capital of United States dollars ("**USD**") 100,000 divided into 100,000 shares with a par value of USD1.00 each. On the same date, 45,500 shares of the Company with nominal value of USD45,500 (equivalent to approximately RMB293,000) had been issued to the Company's shareholders.

As at August 30, 2023, the authorized share capital of the Company was re-designated and subdivided from US\$100,000 divided into 100,000 Shares with a par value of US\$1.00 each to US\$100,000 divided into 177,485,000 Shares with a par value of US\$0.0005 each and 22,515,000 Series A Preferred Shares with a par value of US\$0.0005 each.

Notes to the Condensed Consolidated Financial Statements

For the six months ended June 30, 2025

16. Share Capital (Continued)

	Number of shares	Par value USD	Share capital USD'000
Authorized			
As at January 1, 2024 (audited)	177,485,000	0.0005	89
Converted from Series A Preferred Shares	22,515,000	0.0005	11
As at December 31, 2024 (audited) and June 30, 2025 (unaudited)	200,000,000	0.0005	100

	Number of shares	Par value USD	Amount USD'000	Equivalent amount of ordinary shares RMB'000
Issued and fully paid				
As at January 1, 2024 (audited)	100,000,000	0.0005	50	322
Reclassification from financial liabilities at FVTPL (note i)	22,515,000	0.0005	11	80
Issue of shares upon IPO (note ii)	34,151,800	0.0005	17	122
As at December 31, 2024 (audited) and June 30, 2025 (unaudited)	156,666,800	0.0005	78	524

Notes:

- i. Upon IPO, each Series A Preferred Share was automatically converted into ordinary shares based on the conversion price of 1:1 without the payment of any additional consideration. The Series A Preferred Shares meet the definition of equity as the Group has no contractual obligation to deliver cash or a variable number of shares. Therefore, the Series A Preferred Shares were reclassified from financial liabilities to equity at their fair value, resulting in an increase of share capital of RMB80,000 and an increase of share premium of RMB276,663,000.
- ii. On May 24, 2024, 34,151,800 ordinary shares with par value of USD0.0005 each were issued at HK\$13.50 by way of IPO, resulting in an increase of the share capital of USD17,000 (equivalent to approximately HK\$134,000 or RMB122,000) and an increase of share premium of approximately HK\$460,915,000 (equivalent to approximately RMB419,655,000).

Notes to the Condensed Consolidated Financial Statements

For the six months ended June 30, 2025

17. Share-Based Payment Transactions

Restricted Share Unit Plan

The purpose of the Employee Share Incentive Plan ("**Restricted Share Unit/RSU Plan**") was to provide incentives to employees and directors in order to promote the success of the business of the Group. To implement the RSU Plan, the Company used employee stock ownership platforms (the "**Shareholding Platforms**"), namely Sunho Stellar Investments Limited which was established in April 2021 to hold the Company issued 4,500 shares, representing 9% of the shares of the Company.

On May 6, 2023, the employees of the Group and director of the Company were granted a total of 3,000 shares of the Shareholding Platforms, representing 6% of the shares of the Company.

1,500 shares transferred to Sunho Wisdom did not attached any condition and were fully vested during the year ended December 31, 2023.

On the assumption of the subdivision of shares, each share of US\$1.00 par value was reconstituted into 2,000 shares of US\$0.0005 par value each.

Details of the RSU Plan were disclosed in the Group's consolidated financial statements for the year ended December 31, 2024.

Set out below are details of the movements of equity-settled share-based transactions during the period:

	As at January 1, 2025 (audited)	Granted during the period (unaudited)	Forfeited during the period (unaudited)	Vested during the period (unaudited)	As at June 30, 2025 (unaudited)
IPO-based RSU	6,000,000	–	–	(1,200,000)	4,800,000
Directors	500,000	–	–	(100,000)	400,000
Employees	5,500,000	–	–	(1,100,000)	4,400,000
	6,000,000	–	–	(1,200,000)	4,800,000
Weighted average exercise price (USD)	–	–	–	–	–

For the six months ended June 30, 2025, the Group recognized further RSU-related expenses of RMB7,527,000 (the six months ended June 30, 2024: RMB19,483,000).

Notes to the Condensed Consolidated Financial Statements

For the six months ended June 30, 2025

18. Fair Value Measurements of Financial Instruments

Fair value measurements and valuation processes

Some of the Group's financial instruments are measured at fair value at the end of each reporting period. The following table gives information about how the fair values of these financial liabilities are determined (in particular, the valuation technique(s) and inputs used), as well as the level of the fair value hierarchy into which the fair value measurements are categorised (Levels 1 to 3) based on the degree to which the inputs to the fair value measurements is observable.

- Level 1 fair value measurements are based on quoted prices (unadjusted) in active market for identical assets or liabilities;
- Level 2 fair value measurements are those derived from inputs other than quoted prices included within Level 1 that are observable for the asset or liability, either directly (i.e. as prices) or indirectly (i.e. derived from prices); and
- Level 3 fair value measurements are those derived from valuation techniques that include inputs for the asset or liability that are not based on observable market data (unobservable inputs).

Fair value of the Group's financial assets and financial liabilities that are measured at fair value on a recurring basis

Financial assets/liabilities	Fair value as at		Fair value hierarchy	Valuation technique(s) and key input(s)
	June 30, 2025 RMB'000 (unaudited)	December 31, 2024 RMB'000 (audited)		
Unlisted equity investment at FVTOCI	910	910	Level 2	Quoted recent transaction price
Wealth management products at FVTPL	—	158,825	Level 2	Net asset value quoted by financial institutions

The directors consider that the carrying amounts of financial assets and financial liabilities measured at amortized cost in the condensed consolidated financial statements approximate their respective fair values at the end of the reporting period.

Notes to the Condensed Consolidated Financial Statements

For the six months ended June 30, 2025

19. Related Party Transactions

(a) Compensation of key management personnel

The emoluments of key management for the six months ended June 30, 2025 are as follows:

	Six months ended June 30,	
	2025	2024
	RMB'000	RMB'000
	(unaudited)	(unaudited)
Salaries and other benefits	1,168	1,167
Discretionary bonus (note)	116	60
Retirement benefit scheme contributions	34	33
Share-based payments	7,527	19,843
	8,845	21,103

Note: Discretionary bonus is determined based on their duties and responsibilities of the relevant individuals within the Group and the Group's performance.

20. Capital Commitment

	As at	As at
	June 30, 2025	December 31, 2024
	RMB'000	RMB'000
	(unaudited)	(audited)
Capital expenditure contracted for but not provided in the consolidated financial statements:		
— Leasehold land and equipment	19,125	23,333

Definitions

In this report, unless the context otherwise requires, the following words and expressions shall have the following meanings.

"associate(s)"	has the meaning ascribed to it under the Listing Rules
"Audit Committee"	the audit committee of the Board
"Board"	the board of Directors of the Company
"BVI"	the British Virgin Islands
"Company", "our Company" or "we"	Sunho Biologics, Inc. (盛禾生物控股有限公司), an exempted company with limited liability incorporated in the Cayman Islands on May 14, 2021 and the issued Shares of which are listed on the Stock Exchange (Stock Code: 2898)
"Controlling Shareholders"	has the meaning ascribed to it under the Listing Rules and unless the context otherwise requires, refers to Mr. Zhang Feng, Sunho Fortune, Innovalue Investments, Sunho Wisdom, No5XJR and Sunho Stellar
"Core Products"	namely, IAH0968, IAP0971 and IAE0972
"Director(s)"	the director(s) of our Company
"FDA"	U.S. Food and Drug Administration
"FVTPL"	fair value through profit or loss
"GMP"	good manufacturing practice, a system for ensuring that products are consistently produced and controlled according to quality standards, which is designed to minimize the risks involved in any pharmaceutical production that cannot be eliminated through testing the final product. It is also the practice required in order to conform to the guidelines recommended by agencies that control the authorization and licensing of the manufacture and sale of pharmaceutical products
"Group"	collectively, the Company and its subsidiaries
"HK\$"	Hong Kong dollars, the lawful currency of Hong Kong
"Hong Kong"	the Hong Kong Special Administrative Region of the PRC
"IND"	investigational new drug or investigational new drug application, also known as clinical trial application, or CTA, in China

Definitions

"Innovalue Investments"	Innovalue Investments Limited, a company incorporated in the BVI with limited liability on April 8, 2021 and one of the Controlling Shareholders
"Listing Date"	May 24, 2024
"Listing Rules"	the Rules Governing the Listing of Securities on The Stock Exchange of Hong Kong Limited, as amended, supplemented or otherwise modified from time to time
"Model Code"	Model Code for Securities Transactions by Directors of Listed Issuers as set out in Appendix C3 to the Listing Rules
"NMPA"	the National Medical Products Administration of PRC
"No5XJR"	No5XJR Limited, a company incorporated in the BVI with limited liability on April 14, 2021 and one of the Controlling Shareholders
"PRC" or "China" or "Mainland China"	the People's Republic of China, which for the purpose of this report, excludes Hong Kong, the Macau Special Administrative Region of the PRC and Taiwan
"Prospectus"	prospectus of the Company dated May 16, 2024
"R&D"	research and development
"Reporting Period"	the six months ended June 30, 2025
"RMB"	Renminbi, the lawful currency of the PRC
"RSU"	restricted share unit
"RSU Scheme"	the RSU scheme approved and adopted by the Company on August 2, 2023
"SFO"	the Securities and Futures Ordinance (Chapter 571 of the Laws of Hong Kong), as amended, supplemented or otherwise modified from time to time
"Share(s)"	ordinary share(s) in the share capital of our Company with a par value of USD0.0005 each
"Shareholder(s)"	holder(s) of our Share(s)
"Stock Exchange"	The Stock Exchange of Hong Kong Limited

Definitions

"subsidiary(ies)"	has the meaning ascribed to it under the Listing Rules
"Sunho Fortune"	Sunho Fortune Investments Limited, a company incorporated in the BVI with limited liability on April 9, 2021 and one of the Controlling Shareholders
"Sunho Stellar"	Sunho Stellar Investments Limited, a company incorporated in the BVI with limited liability on April 9, 2021, our share incentive platform and one of the Controlling Shareholders
"Sunho Wisdom"	Sunho Wisdom Investments Limited, a company incorporated in the BVI with limited liability on April 14, 2021 and one of the Controlling Shareholders
"treasury share(s)"	has the meaning ascribed to it under the Listing Rules
"U.S."	the United States of America, its territories, its possession and all areas subject to its jurisdiction
"USD"	United States dollars, the lawful currency of the U.S.
"%"	per cent