
**UNITED STATES
SECURITIES AND EXCHANGE COMMISSION
Washington, D.C. 20549**

FORM 6-K

**REPORT OF FOREIGN PRIVATE ISSUER
PURSUANT TO RULE 13a-16 OR 15d-16
UNDER THE SECURITIES EXCHANGE ACT OF 1934**

For the Month of January 2023

Commission File Number: 001-39992

Immunocore Holdings plc

(Translation of registrant's name into English)

**92 Park Drive
Milton Park
Abingdon, Oxfordshire OX14 4RY
United Kingdom
(Address of principal executive office)**

Indicate by check mark whether the registrant files or will file annual reports under cover of Form 20-F or Form 40-F:

Form 20-F Form 40-F

INCORPORATION BY REFERENCE

The information in this Report on Form 6-K (“Report”), other than Exhibit 99.1 hereto, shall be deemed to be incorporated by reference into the registration statements on Form S-8 (File Nos. 333-255182 and 333-265000) and the registration statement on Form F-3ASR (File No. 333-264105) of Immunocore Holdings plc (the “Company”) and to be a part thereof from the date on which this Report is furnished, to the extent not superseded by documents or reports subsequently filed or furnished.

Exhibit 99.1 hereto is being furnished and shall not be deemed “filed” for purposes of Section 18 of the Securities Exchange Act of 1934 (the “Exchange Act”) or otherwise subject to the liabilities of that section, nor shall it be deemed incorporated by reference in any filing under the Securities Act of 1933 or the Exchange Act.

INFORMATION CONTAINED IN THIS REPORT ON FORM 6-K

Presentation at 41st Annual J.P. Morgan Healthcare Conference

On Wednesday, January 11, 2023, Bahija Jallal, the Chief Executive Officer of the Company, presented at the 41st Annual J.P. Morgan Healthcare Conference. The presentation, which was webcasted, is available in the “Investors/Media” section of the Company’s website, located at www.immunocore.com. A copy of the presentation is furnished as Exhibit 99.1 and is incorporated herein by reference.

The key highlights from the presentation include the following:

- The Company has added three new ImmTAC product candidates (targeting PRAME-A24, PRAME-A02-HLE (half-life extended), and PIWIL1) to the Company’s pipeline.
- As of January 2023, the Company has dosed over 500 cancer patients with KIMMTRAK/tebentafusp for the treatment of metastatic uveal melanoma.
- As of January 2023, KIMMTRAK is approved in over 30 countries, with continued global commercial expansion planned for 2023-2024.
- The Company is starting a Phase 2/3 clinical trial to investigate the potential of tebentafusp for the treatment of advanced cutaneous melanoma. The Company estimates this expansion opportunity would be a potential addressable patient population that is 2-4 times larger than the opportunity for uveal melanoma.
- The Company expects to report initial data from the monotherapy and combination arms of the Phase 1/2 dose escalation trial of IMC-F106C (PRAME-A02) by the first half of 2024.
- The Company believes IMC-R117C is the first PIWIL1 targeted immunotherapy and plans to submit an investigational new drug application in the fourth quarter of 2023.
- The Company plans to report data from the single ascending dose portion of the Phase 1 clinical trial of IMC-M113V, the Company’s ImmTAV molecule targeting the human immunodeficiency virus, in 2023.

CAUTIONARY NOTE ON FORWARD-LOOKING STATEMENTS

This Report contains forward-looking statements within the meaning of the Private Securities Litigation Reform Act of 1995. Words such as “may,” “will,” “expect,” “plan,” “anticipate,” “estimate,” “intend” and similar expressions (as well as other words or expressions referencing future events, conditions or circumstances) are intended to identify forward-looking statements. These statements include, but are not limited to, statements regarding the Company’s strategic priorities, pipeline and expansion thereof, the therapeutic potential and expected clinical benefits, of the Company’s products and product candidates, and the progress, timing, scope, expansion and results of Immunocore’s existing and planned clinical trials, including statements regarding continued global commercial expansion of KIMMTRAK, the estimated potential addressable patient population for tebentafusp for the treatment of advanced cutaneous melanoma, the timing for reporting initial data from the monotherapy and combination arms of the Phase 1/2 dose escalation trial of IMC-F106C (PRAME-A02), the Company’s belief that IMCR-R117C is the first PIWIL1 targeted immunotherapy, the timing for submission of an investigational new drug application for IMC-R117C, and the timing for reporting data from the single ascending dose portion of the Phase 1 clinical trial of IMC-M113V. These forward-looking statements are based on the Company’s expectations and assumptions as of the date of this Report, and are subject to a number of risks and uncertainties that could cause actual results to differ materially and adversely from those set forth in or implied by such forward-looking statements, many of which are beyond the Company’s control. Actual results may differ materially from those expressed or implied by these forward-looking statements. For a discussion of risk factors that may cause the Company’s actual results to differ from those expressed or implied in the forward-looking statements in this Report, you should refer to the Company’s filings with the U.S. SEC, including the “Risk Factors” sections contained therein. Except as required by law, the Company undertakes no obligation to publicly update any forward-looking statements, whether as a result of new information, future events or otherwise. You should, therefore, not rely on these forward-looking statements as representing the Company’s views as of any date subsequent to the date of this Report.

EXHIBIT INDEX

Exhibit No.	Description
99.1	Annual J.P. Morgan Healthcare Conference presentation, dated January 11, 2023.

SIGNATURES

Pursuant to the requirements of the Securities Exchange Act of 1934, the registrant has duly caused this report to be signed on its behalf by the undersigned, thereunto duly authorized.

IMMUNOCORE HOLDINGS PLC

Date: January 11, 2023

By: /s/ Bahija Jallal, Ph.D.

Name: Bahija Jallal, Ph.D.

Title: Chief Executive Officer



IMMUNOCORE

Transformative Medicines for Patients

Bahija Jallal, PhD – Chief Executive Officer
41st Annual J.P. Morgan Healthcare Conference

JANUARY 11TH, 2023

Forward-looking statement

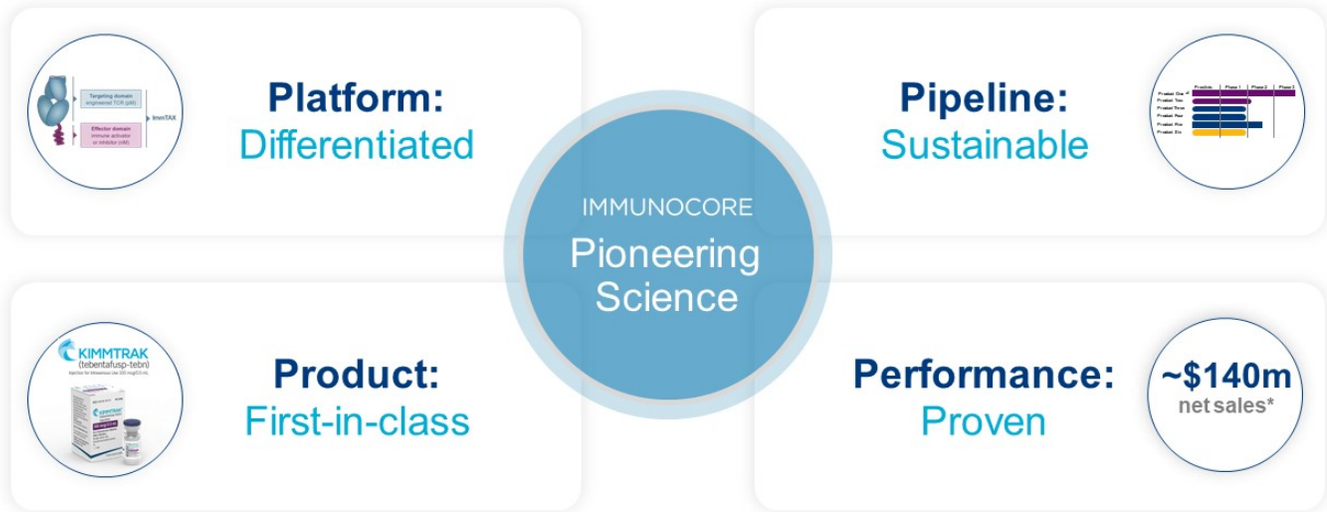
This presentation contains forward-looking statements within the meaning of the Private Securities Litigation Reform Act of 1995. Words such as “may,” “can,” “will,” “believe,” “expect,” “plan,” “anticipate,” “potential” and similar expressions (as well as other words or expressions referencing future events or circumstances) are intended to identify forward-looking statements. All statements, other than statements of historical facts, included in this presentation are forward-looking statements. These statements include, but are not limited to, statements regarding the marketing, therapeutic potential, and expected clinical benefits, including extended overall survival benefit and reduction in circulating tumor DNA, of Immunocore’s products and product candidates; expectations regarding the development of Immunocore’s pipeline and the design, progress, timing, enrollment, scope, expansion and results of Immunocore’s existing, planned and other future clinical trials and IND enabling studies, including the targeted delivery of IND for three new product candidates, the expansion of, and timing for reporting data from the monotherapy and combination arms of, the PRAME-A02 trial and the initiation of the multiple ascending dose portion of, and timing for reporting data from the single ascending dose portion of, the IMC-M113V Phase 1 HIV clinical trial; the ability of TCR therapeutics to target approximately 90% of the human proteome; statements regarding the durability, efficacy and toleration of Immunocore’s product candidates; expectations regarding the commercialization of KIMMTRAK, including potential growth opportunities and trends and increasing access to KIMMTRAK; expectations regarding the value proposition of KIMMTRAK in metastatic uveal melanoma (mUM) and advanced melanoma; expectations regarding the potential market size and opportunity for Immunocore’s products and product candidates, including statements with respect to potential patient population; expectations regarding the number of patients that PRAME-A02 has the potential to benefit; statements that IMC-R117C is a first-in-class PIVL1-targeted immunotherapy under development; statements regarding the planned IND timing for IMC-R117C; expectations regarding future milestones; future development plans of tebentafusp and Immunocore’s other product candidates; the ability to obtain and maintain regulatory approval for its products and product candidates; expectations regarding the sustained or potential commercial performance and uptake of KIMMTRAK and Immunocore’s other product candidates, if approved; expectations regarding Immunocore’s management of resources and expected cash runway; and preliminary unaudited net sales and cash and cash equivalents of KIMMTRAK and tebentafusp; and the validation of the ImmTAC platform. These forward-looking statements are based on management’s current expectations and beliefs and are subject to a number of risks, uncertainties and important factors that may cause actual events or results to differ materially and adversely from those expressed or implied by any forward-looking statements, many of which are beyond Immunocore’s control. These include, without limitation, risks and uncertainties related to the impact of worsening macroeconomic conditions and the ongoing and evolving COVID-19 pandemic, the war in Ukraine or global geopolitical tension on Immunocore’s business, strategy, clinical trials, financial position and anticipated milestones, including Immunocore’s ability to conduct ongoing and planned clinical trials; Immunocore’s ability to obtain and maintain regulatory approval of its product candidates; Immunocore’s ability to obtain clinical supply of current or future product candidates or commercial supply of KIMMTRAK or any future approved products, including as a result of supply chain disruptions; Immunocore’s ability to develop, manufacture and commercialize its product candidates; Immunocore’s ability and plans to launch, market and sell KIMMTRAK or any future approved products, to continue to establish and expand a commercial infrastructure; Immunocore’s ability to successfully expand the approved indications for KIMMTRAK, or obtain marketing approval for KIMMTRAK in additional geographies in the future; the delay of any current or planned clinical trials, whether due to the COVID-19 pandemic, patient enrollment delays or otherwise; unexpected safety or efficacy data observed during preclinical studies or clinical trials and Immunocore’s ability to successfully demonstrate the safety and efficacy of its product candidates and gain approval of its product candidates on a timely basis, if at all; competition with respect to market opportunities; actions of regulatory agencies, which may affect the initiation, timing and progress of clinical trials or future regulatory approval; Immunocore’s ability to obtain, maintain and enforce intellectual property protection for KIMMTRAK or any product candidates it is developing; clinical trial site activation or enrollment rates that are lower than expected; Immunocore’s need for and ability to obtain additional funding on favorable terms or at all, including as a result of worsening macroeconomic conditions such as rising inflation and interest rates; volatility in the capital markets and related market uncertainty; and the success of Immunocore’s current and future collaborations, partnerships or licensing arrangements. These and other risks and uncertainties are described in greater detail in the section titled “Risk Factors,” in Immunocore’s filings with the Securities and Exchange Commission, including Immunocore’s most recent Annual Report on Form 20-F, as supplemented by its most recent filings that Immunocore has made or may make with the SEC in the future. Such risks may be amplified by the COVID-19 pandemic and its potential impact on Immunocore’s business and the overall global economy. Any forward-looking statements represent Immunocore’s views only as of the date of this presentation and should not be relied upon as representing its views as of any subsequent date. Immunocore does not assume any obligation to update any forward-looking statements, except as may be required by law. In addition, as the reported net sales and cash and cash equivalents in this presentation are preliminary, have not been audited and are subject to change pending completion of our audited financial statements for the year ended December 31, 2022, it is possible that Immunocore or its independent registered public accounting firm may identify items that require Immunocore to make adjustments to the amount included in this presentation, and such changes could be material. Additional information and disclosures would also be required for a more complete understanding of Immunocore’s financial position and results of operations as of December 31, 2022.

Certain information contained in this presentation relates to or is based on studies, publications, surveys, and other data obtained from third-party sources and Immunocore’s own internal estimates and research. While Immunocore believes these third-party sources to be reliable as of the date of this presentation, it has not independently verified, and makes no representation as to the adequacy, fairness, accuracy, or completeness of, any information obtained from third-party sources.

KIMMTRAK™ is a trademark owned or licensed to Immunocore.

Building a fully integrated sustainable biotechnology company

Track record from research to commercialization



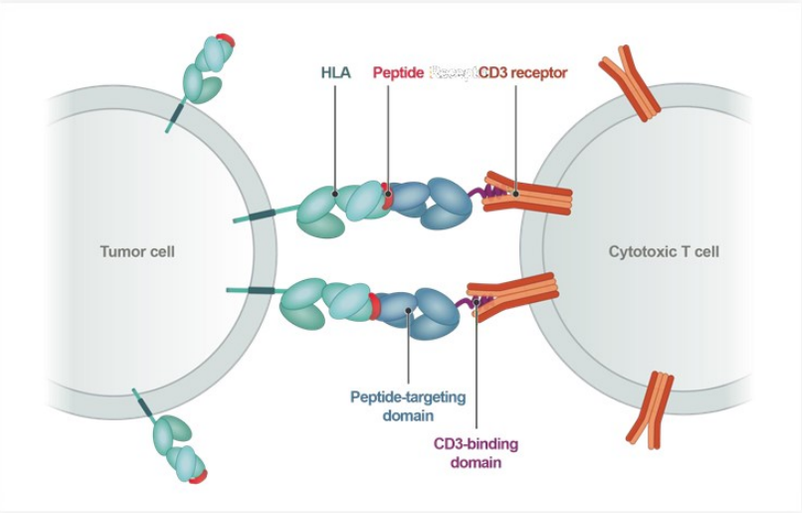
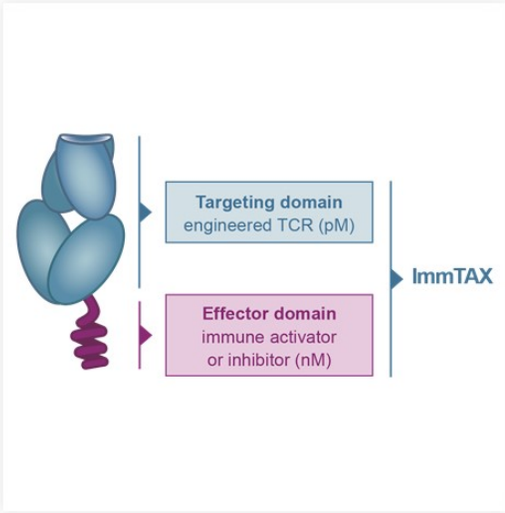
* Net sales* refers to total net product and pre-product revenue of KIMMTRAK and tebentafusp based on December 31, 2022 convenience rate of of £1 to \$1.21. Preliminary net sales are approximated and unaudited.

Our mission

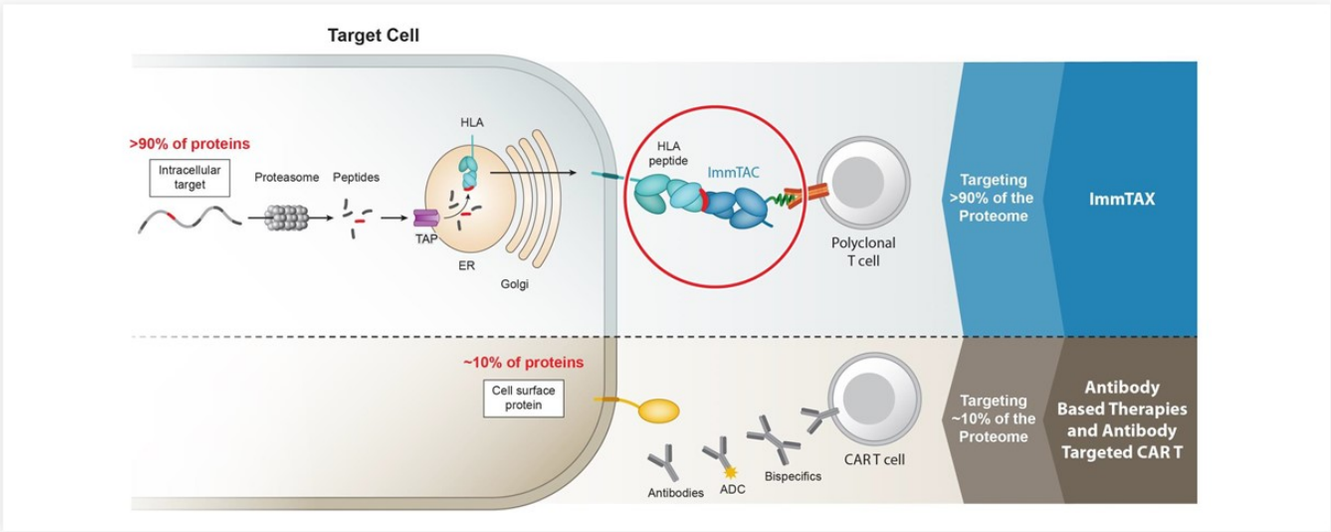


To **radically improve** outcomes **for patients with cancer, infectious diseases, and autoimmune conditions** by pioneering and delivering **transformative** medicines

Harnessing the immune system to fight disease with targeted, off-the-shelf, bispecific, soluble T cell receptors (TCRs)



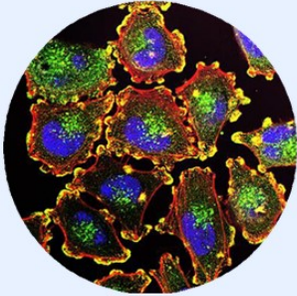
TCR therapeutics target >90% of the human proteome



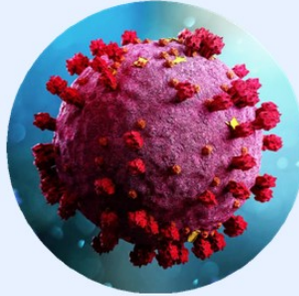
Our platform is modular

Applicable across 3 therapeutic areas

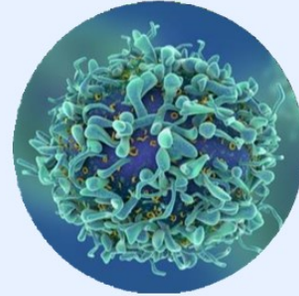
Oncology



Infectious Diseases



Autoimmune Conditions



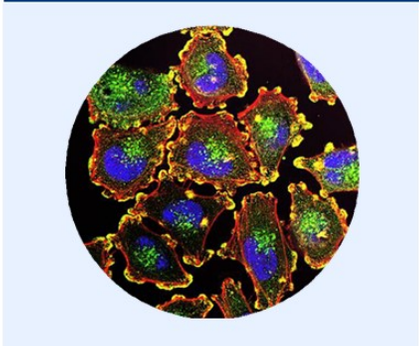
UPREGULATION
OF THE IMMUNE SYSTEM



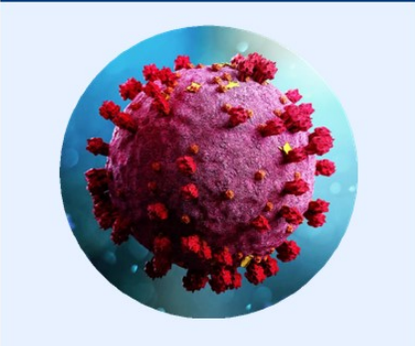
DOWNREGULATION
OF THE IMMUNE SYSTEM

Today's presentation will cover oncology and infectious diseases

Oncology



Infectious Diseases



Autoimmune Conditions



UPREGULATION
OF THE IMMUNE SYSTEM

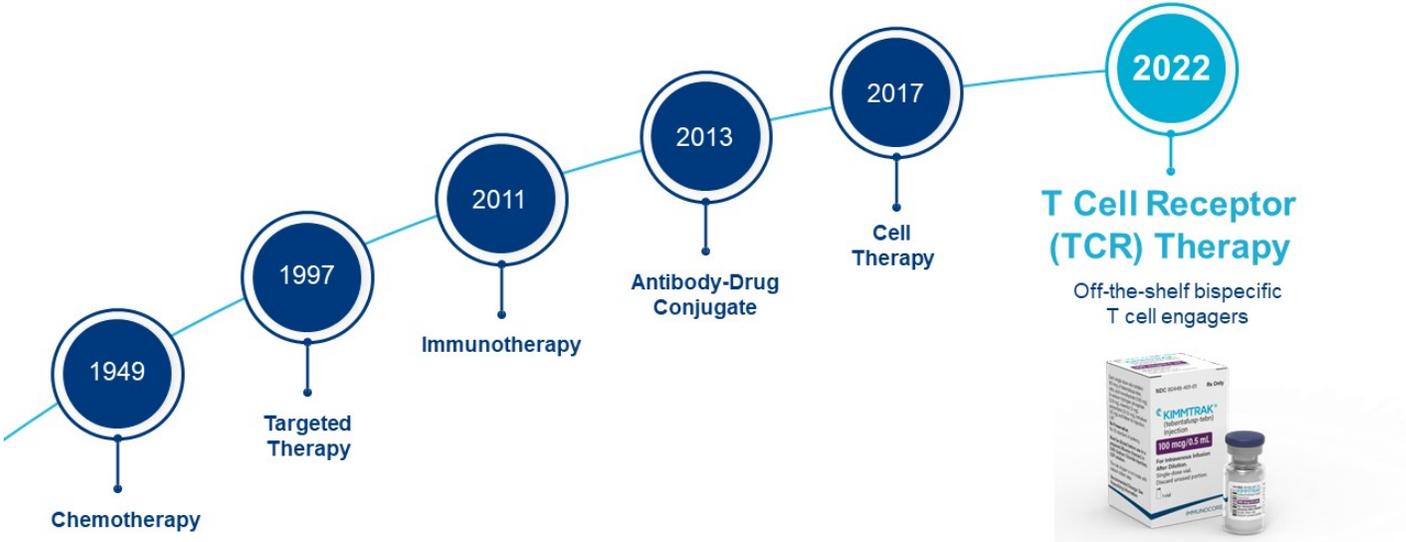


DOWNREGULATION
OF THE IMMUNE SYSTEM

The next chapter in oncology



We have written the next chapter in cancer treatment





KIMMTRAK®

We are leading the way in TCR therapeutics

1st
KIMMTRAK®
(tebentafusp-tebn):
first approved TCR
therapeutic

1st
First and only
FDA-approved treatment
for metastatic uveal
melanoma

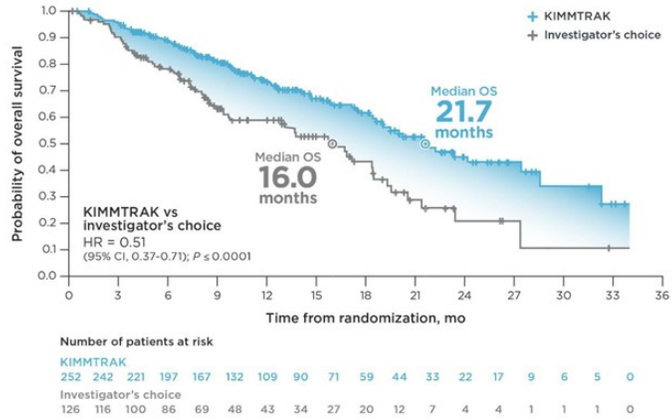
1st
First T-cell engager
to show Overall
Survival (OS) in
solid tumor

KIMMTRAK prolongs overall survival (Hazard ratio: 0.51)

First-in-class, off-the-shelf, bispecific TCR with median OS of 21.7 months



Overall Survival benefit in patients treated with KIMMTRAK or investigator's choice in first-line



Executing on the 2022 global commercial launch of KIMMTRAK

~\$50M

Preliminary Q4 net sales

~\$140M

Preliminary year end net sales

50%

of patients are now first line (1L)

30+

Country approvals*

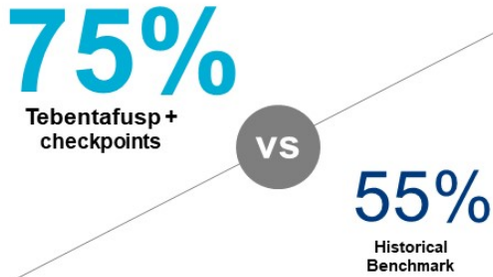


>500 patients treated with KIMMTRAK since Phase 3 data

1. "Net sales" refers to total net product and pre-product revenue of KIMMTRAK and tebentafusp based on December 31, 2022 convenience rate of of £1 to \$1.21. Preliminary net sales are approximated and unaudited.; 2. including US, UK, Canada, Australia, and countries w/in the EU

Expanding beyond UM to previously-treated cutaneous melanoma

Tebentafusp 1-year OS¹ in CM higher than historical benchmark



Randomized Phase 2/3 with OS endpoint

- ▶ HLA*A2:01
- ▶ Cutaneous melanoma
- ▶ Prior anti-PD1
 - Progression within 6 months last dose
- ▶ Prior ipilimumab
- ▶ Prior TKI (if BRAFm)



Primary endpoint: OS

2-4X larger potential addressable patient population than uveal melanoma

A photograph showing a doctor in a white coat and glasses, with a stethoscope, sitting and holding a tablet. An elderly woman with short grey hair, wearing a light-colored knitted cardigan over a tan top and blue jeans, is sitting next to him, looking at the tablet with a thoughtful expression. The background is a bright, indoor setting, possibly a home or a private care facility, with a yellow chair visible on the right. The entire image is overlaid with a semi-transparent blue filter.

PRAME
Franchise:
A02, A24, A02-HLE

PRAME-A02 has the potential to benefit ~150k patients annually

Prevalence of PRAME expression ¹	Tumor type	HLA*02:01+, PRAME+ metastatic patients (G7) ²
70-100%	Endometrial	>10K
	Melanoma	>10K
	Ovarian	>15K
	NSCLC-squamous	>30K
50-70%	NSCLC-adeno	>40K
	SCLC	>15K
	TNBC	>5K
20-50%	SCCHN	>30K
	Gastric	
	RCC	
	Esophageal	
	Cholangiocarcinoma	
	Cervical	

- **Expressed across multiple solid tumors**

- **Negative prognostic marker**

- **Broad and homogeneous expression within key tumors**

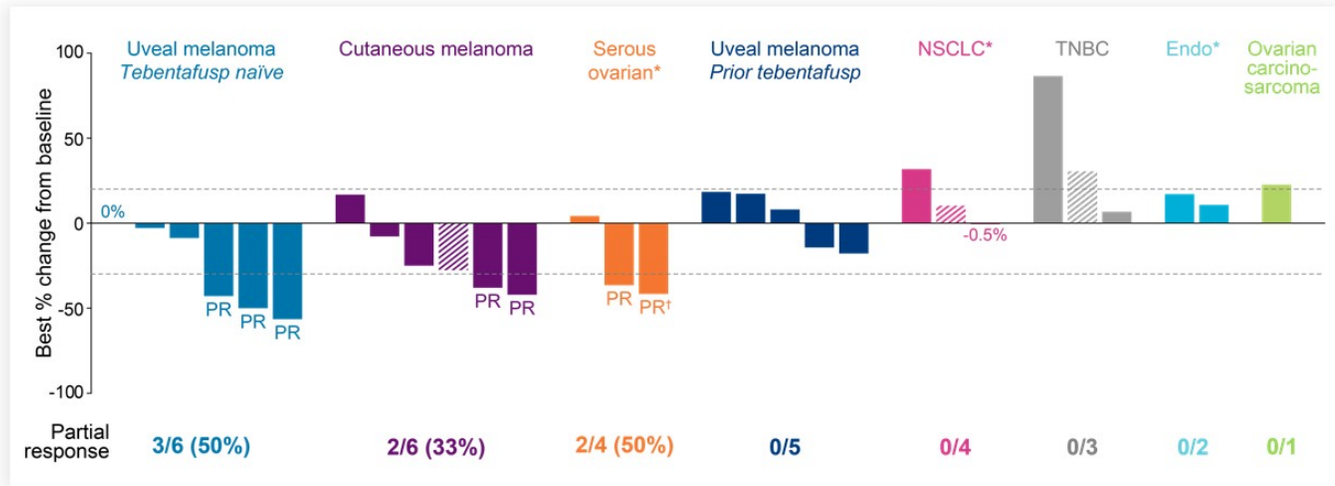
1. PRAME prevalence derived from immunohistochemistry and RTqPCR of patient samples and analysis of TCGA
 2. Epidemiology data from cancer registries and Decision Resources, Annual incidence of metastatic patients

Responses observed in multiple tumor types

IMC-F106C ESMO 2022

PRAME expression[‡]

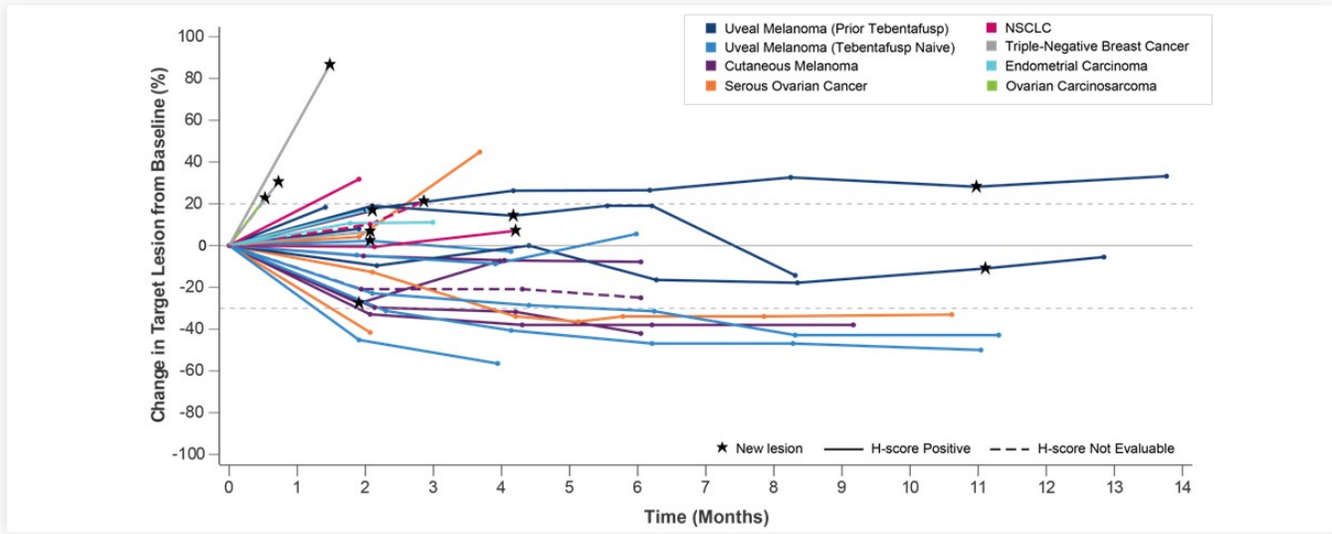
■ Positive ▨ Not evaluable



* Two patients (1 with NSCLC, 1 serous ovarian) discontinued treatment due to PD with scan data not available at DCO; † This serous ovarian patient (H-score 39) had an unconfirmed partial response (uPR) at the time of the ESMO Congress September 2022 presentation, that was subsequently confirmed; ‡ PRAME expression assessed by IHC H-score; Two PRAME-negative patients both had PD (not shown); Endo, endometrial carcinoma; NSCLC, non small cell lung carcinoma; TNBC, triple-negative breast cancer; Hamid, O., et al. *Annals of Oncology* (ESMO 2022) 33 (suppl_7): S331-S355.

Majority of patients have durable tumor response or stabilization

IMC-F106C ESMO 2022

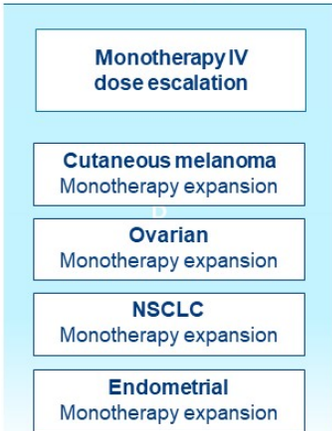


NSCLC, non small cell lung carcinoma
Hamid, O., et al, Annals of Oncology (ESMO 2022) 33 (suppl_7): S331-S355.

Enrolling patients globally in adaptive trial with multiple arms

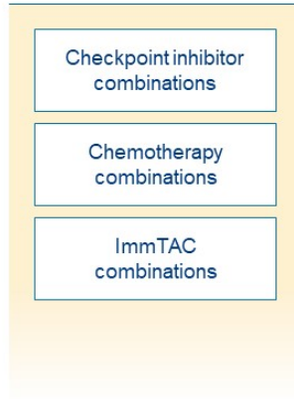
Expanding clinical trial footprint | Aim to understand breadth of clinical activity in solid tumors

Monotherapy



Adaptive design enables flexible expansion size

Combinations



Enables future randomized trials into earlier lines of therapy

Monotherapy activity provides optionality to develop in single arm and randomized trials

Expansion of ImmTAC franchise targeting PRAME

Building on enthusiasm for IMC-F106C targeting PRAME HLA-A02

	Target	HLA subtype	Format	
IMC-F106C	PRAME	HLA-A02	TCRxCD3	<ul style="list-style-type: none">▶ Clinically validated▶ Focus on expanding clinical program
IMC-T119C	PRAME	HLA-A24	TCRxCD3	<ul style="list-style-type: none">▶ Expands potential addressable population by ~30% (G7)▶ High prevalence in Japan
IMC-P115C	PRAME	HLA-A02	TCRxCD3 HLE	<ul style="list-style-type: none">▶ Half-life extended (HLE) for less frequent dosing

HLE, Half-life extension

IMMUNOCORE |



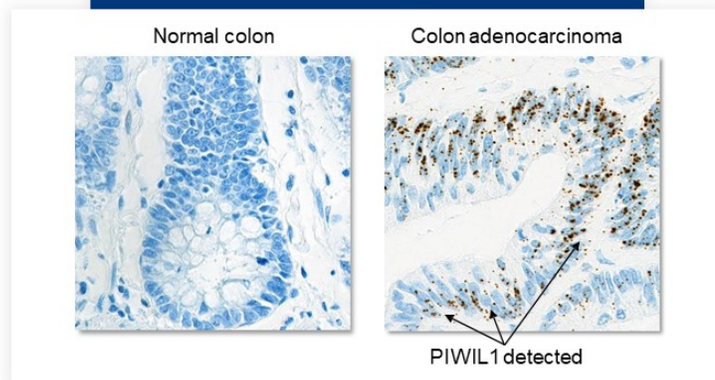
Novel ImmTAC
Candidate for GI
cancers from our
discovery engine

IMC-R117C: A first-in-class immunotherapy targeting PIWIL1

IND planned Q4 2023

- **Negative prognostic marker in multiple cancers**, role in tumor progression
- **Expressed in CRC, historically insensitive to IO**, and across major subgroups[^]
- **25% CRC have broad PIWIL1 expression** (e.g., > 75% of tumor cells positive)

PIWIL1 RNA *in situ* hybridization



Total >35,000* patients/year
positive for PIWIL1 and HLA-A02

PIWIL1, piwi-like protein 1; MSS, microsatellite stable; MSI, Microsatellite instability; CRC, colorectal
* Estimated across colorectal, esophageal, gastric, pancreatic, ovarian, endometroid cancers



Pursuing a
functional cure
in infectious
diseases

Pursuing a functional cure in HBV & HIV

IMC-I109V

HBV Phase 1

- ▶ First cohort (0.8 mcg) reported
- ▶ HBsAg transiently decreased¹
- ▶ Decreases coincided with transient ALT elevations¹

Initial Phase 1 SAD data presented at [EASL 2022 Congress](#)



IMC-M113V

HIV Phase 1

- ▶ Finished single dose escalation and starting Multiple Ascending Dose
- ▶ Initial funding by the Bill & Melinda Gates Foundation²

Phase 1 Single Ascending Dose (SAD) data expected in 2023

1. Bourgeois, et. al. EASL 2022; 2. Program is wholly owned, development costs previously provided by the Bill & Melinda Gates Foundation (BMGF), Immunocore retains all development and commercialization rights in the developed world.

A photograph of two medical professionals, likely doctors, shaking hands. One is wearing a blue lab coat and the other a white lab coat. A stethoscope is visible around the neck of the person in the white coat. The image is partially obscured by a dark blue overlay on the left side where the text is located.

Delivering on our
promise –
Consistent
execution

Delivering leading bispecific TCR pipeline

Multiple candidates in oncology and infectious diseases

Candidate	Target	Indication	IND-enabling	Phase 1	Phase 2	Phase 3	Approved
KIMMTRAK Tebentafusp	gp100	Uveal melanoma		[Progress bar]			
		Advanced melanoma		[Progress bar]			
IMC-F106C	PRAME-A02	Multiple solid tumors		[Progress bar: Monotherapy dose exploration]			
		Multiple solid tumors		[Progress bar: Combinations w/ standards of care]			
		2L+ cutaneous melanoma		[Progress bar]			
		PRR Ovarian*		[Progress bar]			
		Advanced endometrial		[Progress bar]			
		2L+ NSCLC		[Progress bar]			
IMC-P115C	★ PRAME-A02-HLE	Multiple solid tumors		[Progress bar]			
IMC-T119C	★ PRAME-A24	Multiple solid tumors		[Progress bar]			
IMC-R117C	★ PIWIL1	Colorectal, gastric, pancreatic		[Progress bar]			
IMC-C103C ¹	MAGE-A4	Multiple solid tumors		[Progress bar]			
IMC-I109V IMC-M113V ²	Envelope Gag	Hepatitis B Virus (HBV)		[Progress bar]			
		Human Immunodeficiency Virus (HIV)		[Progress bar]			

ONCOLOGY

INFECTIOUS DISEASES

★ New ImmTAC candidate

1. Developed under a co-development/co-promotion collaboration with Genentech; 2. Program is wholly owned, development costs being provided by the Bill & Melinda Gates Foundation (BMGF), Immunocore retains all development and commercialization rights in the developed world. * Platinum refractory or resistant serous ovarian carcinoma

Preliminary 2022 Financial Results

Cash runway projected into 2026 with anticipated KIMMTRAK revenues

~\$50M

Q4 preliminary net sales of KIMMTRAK / tebentafusp^{1,2}

~\$140M

YE preliminary net sales of KIMMTRAK / tebentafusp^{1,2}

~\$400M

Preliminary cash and cash equivalents as of December 31, 2022²



*Preliminary financial results are approximated and unaudited. 1. "Net sales" refers to total net product and net pre-product revenue of KIMMTRAK and tebentafusp. 2. Dollar amounts based on conversion rate of approximately 1.21.

Looking ahead

Continuing to write the next chapter of cancer and infectious diseases treatment



Sustain and grow 

Global site expansion for **PRAME-A02** trial
(data by 1H 2024)

Deliver IND for **3 new ImmTAC** candidates

HIV Phase 1 SAD data 2023

Continue **responsible management** of resources

Experienced team with deep scientific & commercial expertise



Bahija Jallal

CEO



Brian Di Donato

CFO & Head of Strategy



David Berman

Head of R&D



Mohammed Dar

CMO



Andy Hooker

VP, CMC & Supply Chain



JoAnn Suzich

Head of Research



Mark Moyer

Head of Regulatory



Ralph Torbay

Head of Commercial



Regulatory approval of KIMMTRAK® in unresectable or metastatic uveal melanoma (mUM) in 30+ countries



THANK YOU

Delivering leading bispecific TCR pipeline

Multiple candidates in oncology and infectious diseases

Candidate	Target	Indication	IND-enabling	Phase 1	Phase 2	Phase 3	Approved	
KIMMTRAK Tebentafusp	gp100	Uveal melanoma	[Progress bar: Phase 1 to Phase 3]					
		Advanced melanoma	[Progress bar: Phase 1 to Phase 2]					
IMC-F106C	PRAME-A02	Multiple solid tumors	[Progress bar: Phase 1 to Phase 2] Monotherapy dose exploration					
		Multiple solid tumors	[Progress bar: Phase 1 to Phase 2] Combinations w/ standards of care					
		2L+ cutaneous melanoma	[Progress bar: Phase 1 to Phase 2]					
		PRR Ovarian*	[Progress bar: Phase 1 to Phase 2]					
		Advanced endometrial	[Progress bar: Phase 1 to Phase 2]					
		2L+ NSCLC	[Progress bar: Phase 1 to Phase 2]					
IMC-P115C	★ PRAME-A02-HLE	Multiple solid tumors	[Progress bar: Phase 1 to Phase 2]					
IMC-T119C	★ PRAME-A24	Multiple solid tumors	[Progress bar: Phase 1 to Phase 2]					
IMC-R117C	★ PIWIL1	Colorectal, gastric, pancreatic	[Progress bar: Phase 1 to Phase 2]					
IMC-C103C ¹	MAGE-A4	Multiple solid tumors	[Progress bar: Phase 1 to Phase 2]					
IMC-I109V	Envelope	Hepatitis B Virus (HBV)	[Progress bar: Phase 1 to Phase 2]					
IMC-M113V ²	Gag	Human Immunodeficiency Virus (HIV)	[Progress bar: Phase 1 to Phase 2]					

ONCOLOGY

INFECTIOUS DISEASES

★ New ImmTAC candidate

1. Developed under a co-development/co-promotion collaboration with Genentech; 2. Program is wholly owned, development costs being provided by the Bill & Melinda Gates Foundation (BMGF), Immunocore retains all development and commercialization rights in the developed world. * Platinum refractory or resistant serous ovarian carcinoma