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

FORM 8-K

CURRENT REPORT

Pursuant to Section 13 or 15(d) of the Securities Exchange Act of 1934

Date of Report (Date of earliest event reported): February 25, 2026

CORBUS PHARMACEUTICALS HOLDINGS, INC.

(Exact name of Registrant as Specified in Its Charter)

Delaware
(State or Other Jurisdiction
of Incorporation)

001-37348
(Commission File Number)

46-4348039
(IRS Employer
Identification No.)

500 River Ridge Drive
Norwood, Massachusetts
(Address of Principal Executive Offices)

02062
(Zip Code)

Registrant's Telephone Number, Including Area Code: (617) 963-0100

Not Applicable

(Former Name or Former Address, if Changed Since Last Report)

Check the appropriate box below if the Form 8-K filing is intended to simultaneously satisfy the filing obligation of the registrant under any of the following provisions:

- Written communications pursuant to Rule 425 under the Securities Act (17 CFR 230.425)
- Soliciting material pursuant to Rule 14a-12 under the Exchange Act (17 CFR 240.14a-12)
- Pre-commencement communications pursuant to Rule 14d-2(b) under the Exchange Act (17 CFR 240.14d-2(b))
- Pre-commencement communications pursuant to Rule 13e-4(c) under the Exchange Act (17 CFR 240.13e-4(c))

Securities registered pursuant to Section 12(b) of the Act:

Title of each class	Trading Symbol(s)	Name of each exchange on which registered
Common Stock, par value \$0.0001 per share	CRBP	The Nasdaq Capital Market

Indicate by check mark whether the registrant is an emerging growth company as defined in Rule 405 of the Securities Act of 1933 (§ 230.405 of this chapter) or Rule 12b-2 of the Securities Exchange Act of 1934 (§ 240.12b-2 of this chapter).

Emerging growth company

If an emerging growth company, indicate by check mark if the registrant has elected not to use the extended transition period for complying with any new or revised financial accounting standards provided pursuant to Section 13(a) of the Exchange Act.

Item 7.01 Regulation FD Disclosure.

Corbus Pharmaceuticals Holdings, Inc. updated its presentation used by management to describe its business. A copy of the presentation is furnished as Exhibit 99.1 and incorporated herein by reference.

The information in this Current Report on Form 8-K under Item 7.01, including the information contained in Exhibit 99.1, is being furnished to the Securities and Exchange Commission, and shall not be deemed to be “filed” for the purposes of Section 18 of the Exchange Act, or otherwise subject to the liabilities of that section, and shall not be deemed to be incorporated by reference into any filing under the Securities Act or the Exchange Act, except as shall be expressly set forth by a specific reference in such filing.

Item 9.01 Financial Statements and Exhibits.

(d) The following exhibit is furnished with this report:

Exhibit No.	Description
99.1	Investor Presentation
104	Cover Page Interactive Data File (embedded within the Inline XBRL document).

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.

Corbus Pharmaceuticals Holdings, Inc.

Date: February 25, 2026

By: */s/ Yuval Cohen*

Name: Yuval Cohen

Title: Chief Executive Officer



**Connecting Innovation
to Purpose**

Corporate Presentation

February 25, 2026



www.corbuspharma.com

@corbuspharma

NASDAQ: CRBP

Forward-Looking Statements

This presentation contains certain forward-looking statements within the meaning of Section 27A of the Securities Act of 1933 and Section 21E of the Securities Exchange Act of 1934 and Private Securities Litigation Reform Act, as amended, including those relating to the Company's trial results, product development, clinical and regulatory timelines, market opportunity, competitive position, possible or assumed future results of operations, business strategies, potential growth opportunities, including timing or completion of trials and presentation of data and other statements that are predictive in nature. These forward-looking statements are based on current expectations, estimates, forecasts and projections about the industry and markets in which we operate and management's current beliefs and assumptions. These statements may be identified by the use of forward-looking expressions, including, but not limited to, "expect," "anticipate," "intend," "plan," "believe," "estimate," "potential," "predict," "project," "should," "would" and similar expressions and the negatives of those terms. These statements relate to future events or our financial performance and involve known and unknown risks, uncertainties, and other factors, on our operations, clinical development plans and timelines, which may cause actual results, performance or achievements to be materially different from any future results, performance or achievements expressed or implied by the forward-looking statements. Such factors include those set forth in the Company's filings with the Securities and Exchange Commission. Prospective investors are cautioned not to place undue reliance on such forward-looking statements, which speak only as of the date of this presentation. The Company undertakes no obligation to publicly update any forward-looking statement, whether as a result of new information, future events or otherwise. This presentation includes limited observations derived from separate clinical settings that are not, and should not be interpreted as, direct or indirect head-to-head comparisons of CRB-701, CRB-913 or CRB-601 with any other product. The observations described herein are subject to change as additional data become available, and future clinical trials of CRB-701, CRB-913 or CRB-601 may not reproduce, validate, or otherwise confirm these observations.

All product names, logos, brands and company names are trademarks or registered trademarks of their respective owners. Their use does not imply affiliation or endorsement by these companies.



Key clinical readouts for 2026

CRB-701

Clinical updates in HNSCC (1L combo and 2L+ mono) & cervical (2L+ mono)

CRB-913

12-week dose-range finding data in patients with obesity (n=240)

CRB-601

Dose escalation data

\$173M

Cash, cash equivalents and investments as of November 3, 2025, and approximately 17.6M common shares issued and outstanding (~20.5M fully diluted shares)



CRB-701

Next Generation Nectin-4 Targeting ADC



CRB-701: Re-imagining a Nectin-4 ADC

Safety

Markedly reduced PADCEV[®]-associated toxicities

Convenience

Extend ADC half-life → Reduce dosing frequency

Efficacy

Lower DAR + longer half-life → Dose higher + longer than PADCEV[®]

Strategy

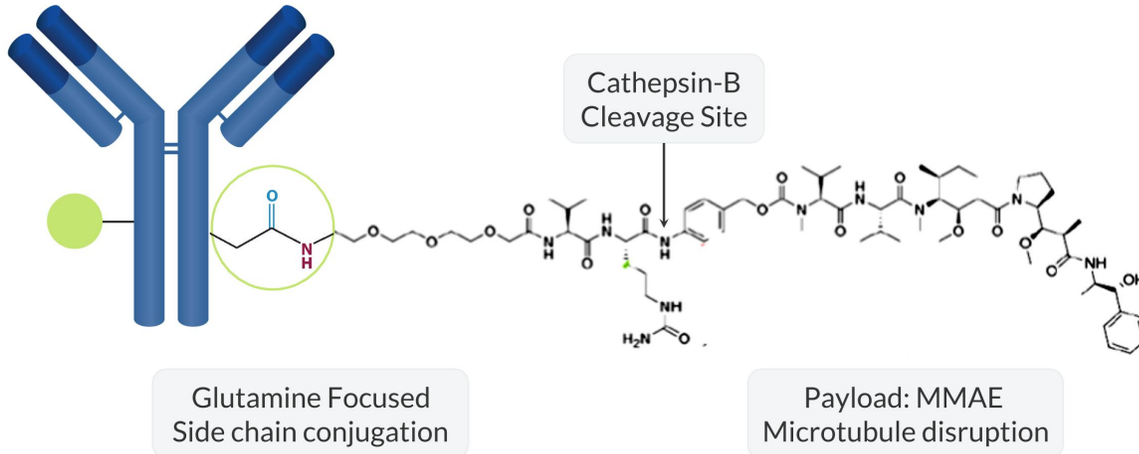
Focus on non-mUC tumors → Avoid competing with PADCEV[®]

CRB-701: Proprietary components → novel design

Novel Nectin-4 Antibody
ADCC + CDC functionality

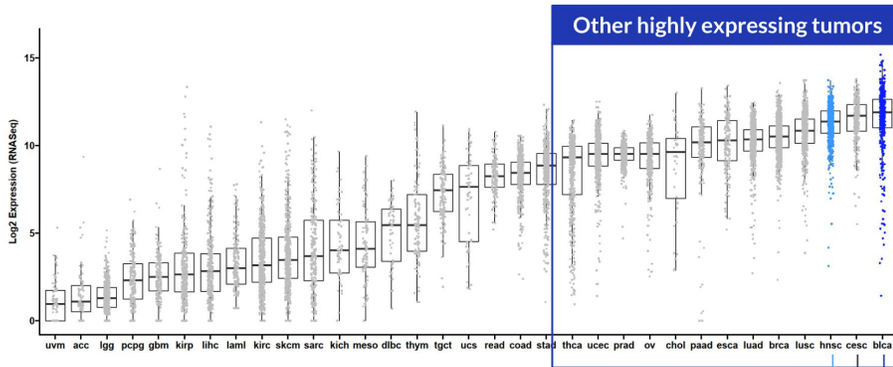
An Improved ADC Construct

- Precise & stable DAR of 2 → Longer half life
- Improved binding affinity & selectivity → 2x rate of internalization vs. PADCEV®
- Improved linker stability → Reduced free MMAE



MMAE = Monomethyl auristatin E. ADCC = antibody-dependent cellular cytotoxicity. CDC = complement dependent cytotoxicity
Source(s): Modified image from Corbus data on file; Corbus data on file

Best responses seen in tumors with highest Nectin-4 expression-mUC, cervical & HNSCC¹

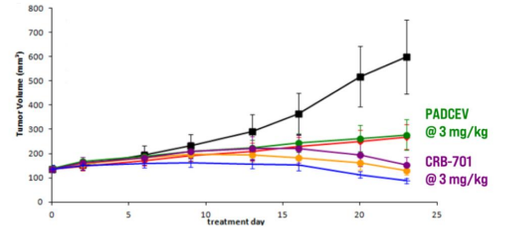


Elevated Nectin-4 expression: urothelial, cervical, head and neck, breast, ovarian, colorectal, rectal, esophageal, gastric, lung, thyroid, prostate, cholangiocarcinoma, pancreatic cancer, testicular cancer

HNSC = Head and neck Cancer (Squamous)
 CESCC = Cervical Cancer (squamous)
 BLCA = Bladder Cancer (urothelial)

CRB-701 demonstrates better efficacy than EV in patient-derived tumor model expressing low levels of Nectin-4²

Mean Tumor Volume ± SEM



- Vehicle
- NS Cisplatin 4 mg/kg
- NS PADCEV 3 mg/kg
- ** CRB-701 2 mg/kg
- *** CRB-701 3 mg/kg
- *** CRB-701 4 mg/kg

CRB-701	PADCEV®
74.5%	53.7%
p<0.05	p=0.70

Key differentiator: Lower levels of free MMAE for CRB-701 vs. PADCEV®

Company	21-day PK	Comparison	% ADC		% Free MMAE	
			C _{max}	AUC _{0-21d}	C _{max}	AUC _{0-21d}
	PADCEV® 1.24 mg/kg Q1W x 3	PADCEV® Benchmark	100%	100%	100%	100%
	2.7 mg/kg Q3W	Matched for MMAE dose (DAR)	183%	274%	35%	38%
	3.6 mg/kg Q3W	2.9-fold PADCEV® ADC Dose®	228%	361%	59%	62%

Source(s):
 PADCEV® reference data from BLA761137 17 December 2019
 Corbus data: ESMO 01 Sep 2025 Data cut

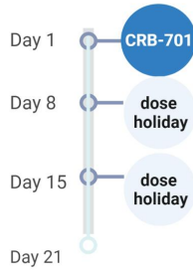
CRB-701: Best-in-class dosing regimen

Clinical Cycle Comparison

Patient / Physician Convenience

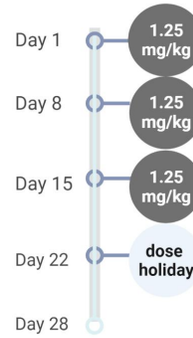
Combination Flexibility

CRB-701

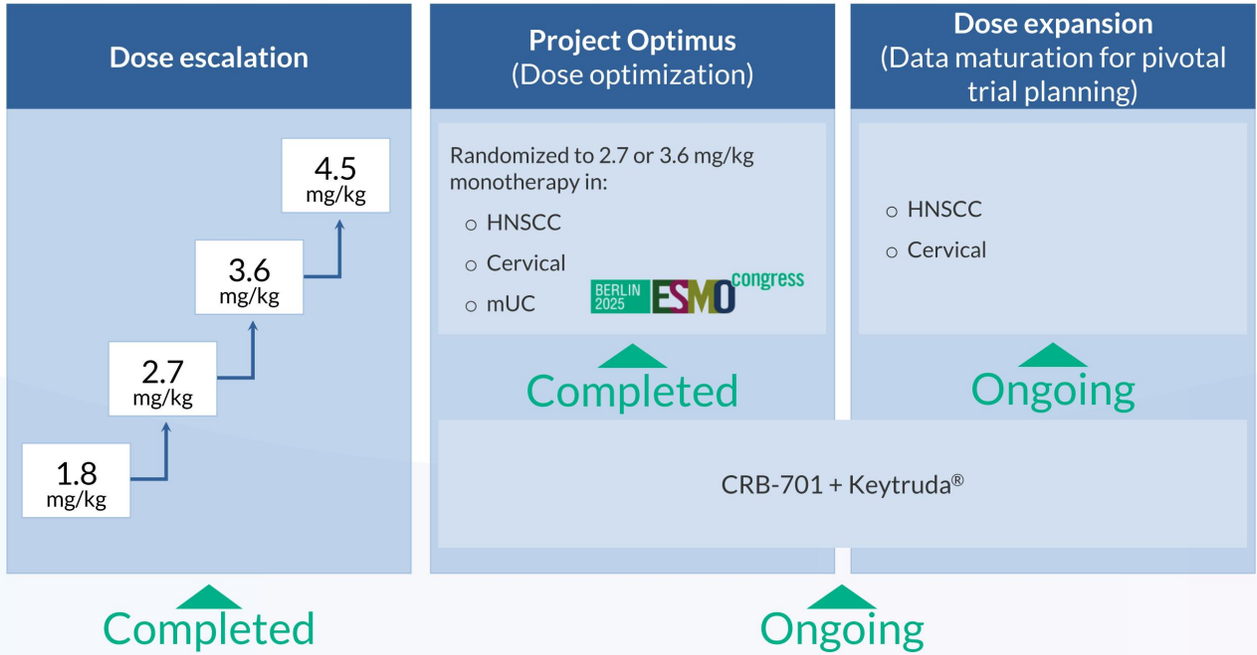


PADCEV[®]

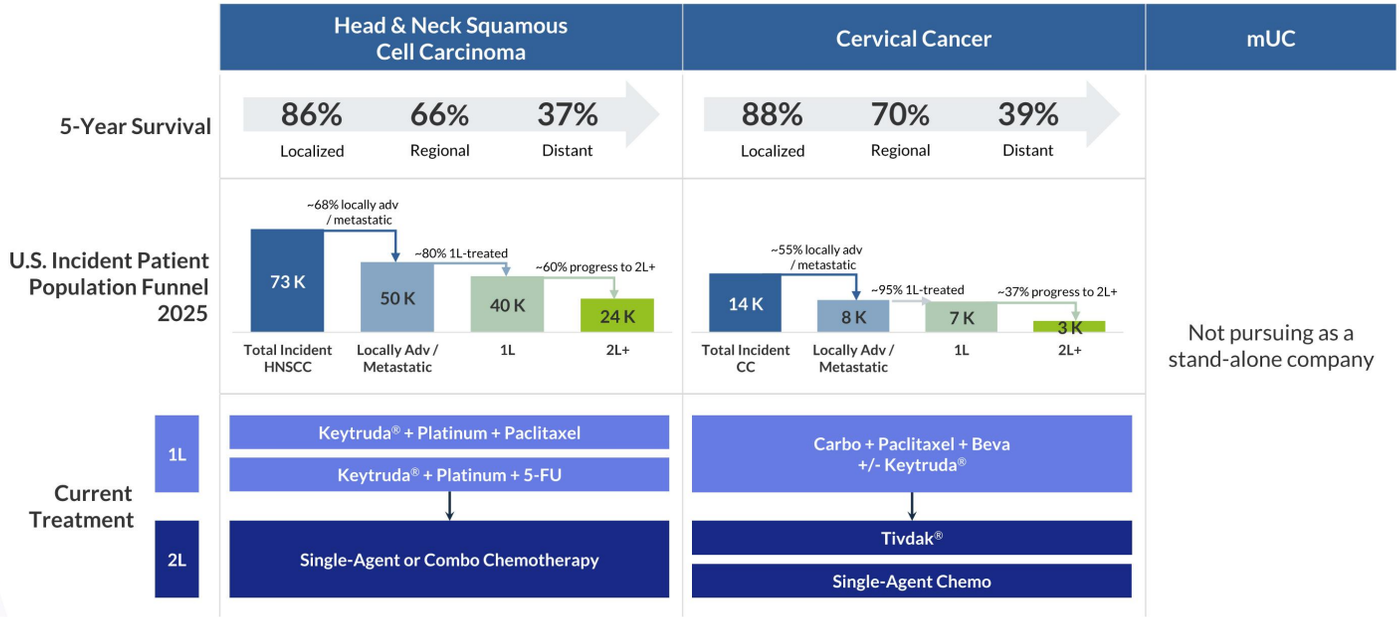
enfortumab vedotin-efjv
Injection for Intravenous Use



CRB-701: Corbus study design (U.S. + Europe)



Emerging indications of interest: HNSCC + cervical cancer



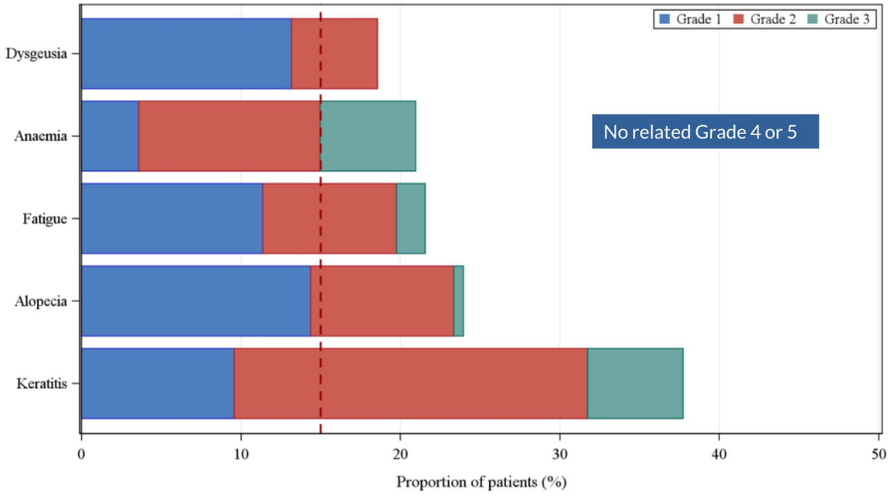
Source: SEER Bladder Cancer; Census.gov; Weir et al., 2021; American Cancer Society; Chu et al., 2022; Hoffman-Censits et al., 2022; SEER Cervical Cancer; Census.gov; Weir et al., 2021; American Cancer Society; Mizuho Analyst Report; Corbus Corporate Deck. SEER Oral Cavity & Pharynx Cancer; SEER Laryngeal Cancer; American Cancer Society; Sanders et al., 2022. LifeSci Consulting Qualitative Market Research



ESMO 2025: Key characteristics & tumor types

Baseline characteristic (as of 9/1/25 data cut)		Enrolled tumor types (n=167)	
Median age (range)	60 (32-90)	HNSCC	60
Sex (M/F)	50.3% / 49.7%	Cervical	54
ECOG PS 0, 1, 2	43.1%, 55.1%, 1.8%	Locally advanced/ mUC	27
Weight in kg mean (range)	72 (32.1-132.8)	NSCLC	7
Prior therapies median (range)	3 (1-9)	TNBC	1
Safety Population	n=167	Endometrial	3
Safety Population dosed with monotherapy CRB-701	n=163	Prostate	1
Efficacy evaluable population (participants with at least 1 post-baseline scans)	n=122	Penile	2
	HNSCC n=41	Ovarian	4
	Cervical n=37	Pancreatic	7
	La/mUC n=23	Missing	1
	Other tumor types n=21		

ESMO 2025: TEAEs ≥15% (n=167)



Adverse Events of Interest	N=167 (%)
Peripheral neuropathy Broad Terms*	8.4%
Eye	
Overall	56.9%
Grade 3	9%
Grade 4 & 5	0
Skin	
Pruritus	14.4%
Dry skin	10.2%
Rash	9.0%
Rash maculo-papular	4.8%
Dermatitis acneiform	3.6%
Erythema	1.8%
Dermatitis bullous	1.2%
Rash pustular	1.2%
Rash erythematous	0.6%
Rash macular	0.6%
Rash pruritic	0.6%
Skin disorder	0.6%
Skin reaction	0.6%
Skin ulcer	0.6%

*Standardized MedDRA Category Search
Sources: ESMO 01 Sep 2025 Data cut



ESMO 2025: Favorable emerging safety profile vs. Nectin-4-MMAE peers



Bicycle



	PADCEV® ¹	BT8009 ²	9MW-2821 ^{3,4}	CRB-701 ⁵	
Upper dose limit	1.25 mg/kg	5 mg/m ²	1.25 mg/kg	2.7mg/kg	3.6mg/kg
Schedule	D1, D8, D15 /28 days	Q1W	D1, D8, D15 /28 days	Q3W	
≥ Grade 3 AE rate	62.5% (n=237/379)	53% (n=24/45)	70%	35.7% (n=25/70)	35.5% (n=27/76)
Peripheral neuropathy (broad terms)	48% (n=182/379)	36% (n=16/45)	22.5% (n=54/240)	8.6% (6/70)	6.6% (5/76)
Rash (broad terms*)	50.7% (n=192/379)	18% (n=8/45)	30% (n=72/240)	32.9% (n=23/70)	23.7% (n=18/76)
Neutropenia (Gr 3)	10% (31/310)	4% (n=2/45)	27.9% (n=67/240)	0%	0%
Dose reduction	27.7% (n=105/379)	27% (n=12/45)	Not released	10% (7/70)	19.7% (15/76)
Dose interruptions	55.9% (n=212/379)	53% (n=24/45)	Not released	38.6% (27/70)	51.3% (39/76)
Discontinuations	20.6% (78/379)	4% (n=2/45)	Not released	5.7% (4/70)	7.9% (6/76)

Source(s):

1. NDA/BLA Multidisciplinary Review and Evaluation BLA 761137 PADCEV® (enfortumab vedotin)
2. Torras, O. Reig, et al. "652P BT8009 monotherapy in enfortumab vedotin (EV)-naïve patients with metastatic urothelial carcinoma (mUC): Updated results of Duravelo-1." *Annals of Oncology* 35 (2024): S515-S516.
3. ASCO 2024, Zhang, et al.
4. SGO plenary March 2024, Yang et al.
5. ESMO 01 Sep 2025 Data cut *Rash (Broad terms): Skin and subcutaneous tissue disorders SOC, excluding alopecia

Safety Summary

Best-in-class for peripheral neuropathy

8.4% (all grade 1 or 2)*

Low rates of skin adverse events

28.7% (excluding alopecia)

Low numbers of Grade ≥ 3 events (3/167**)

Eye toxicities have been manageable with prophylaxis and dose modifications

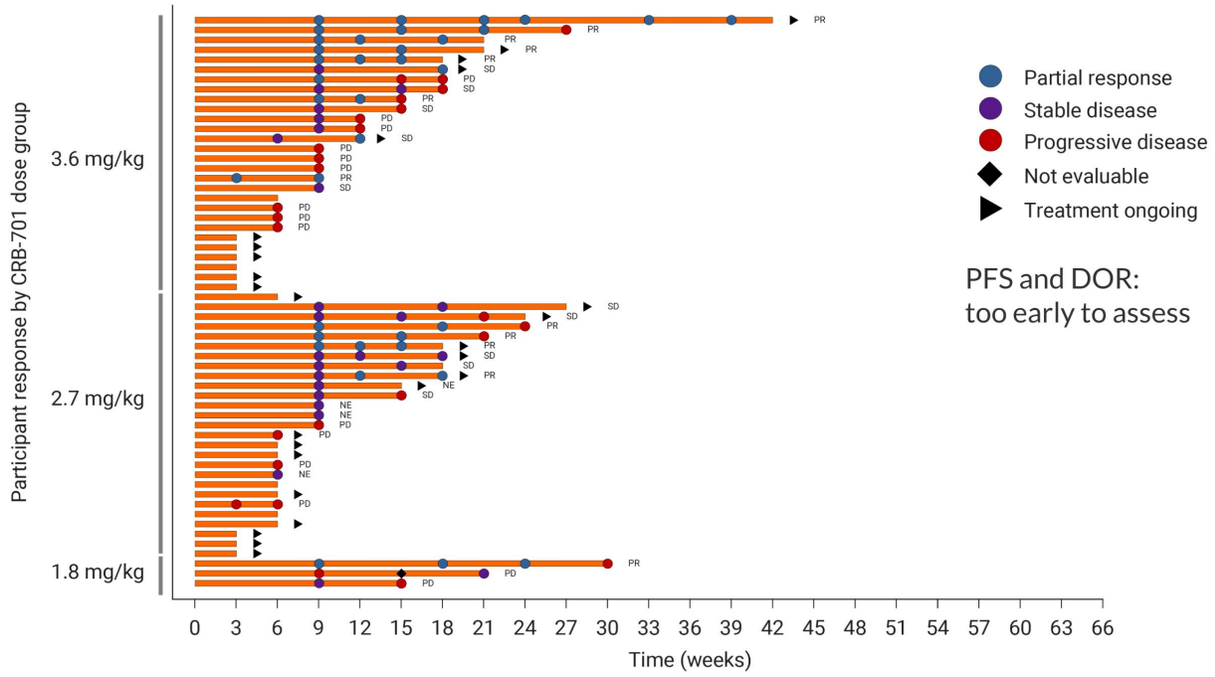
Discontinuations due to eye toxicities have been low (4.2%)

HNSCC baseline characteristics vs. peers

Baseline characteristic	CRB-701*	Petosemtab**	HNSCC PADCEV®***
Median age (range)	62 (35-76)	60 (31-77)	65 (33-81)
Sex (M/F)	90% / 10%	79% / 21%	87% / 13%
ECOG PS 0, 1, 2	48.3%, 50%, 1.7%	30%, 70%, 0%	34.8%, 65.2%, 0%
Prior lines median (range)	3 (1-9)	2 (1-4)	1 line 15.2% 2 lines 17.4% ≥3 lines 67.4%
HPV/P16 Status (Positive/Negative/Missing)	28.3% / 15.0% / 56.7%	46% / 46% / 8%	43.5% / 13% / 43.5%
Disease status at Study Entry (Locally Recurrent/Metastatic)	15% / 85%	Not disclosed	Not disclosed
Nectin-4 H-Score (Range)	13-285	N/A	20-300
PD-L1 Criteria	Agnostic	PD1(L1)-1 Positive	Agnostic

Source(s): * ESMO 01 Sep 2025 Data cut; **ESMO ASIA [data](#) Dec 2024; *** [Swiecicki et al, 2024](#)

ESMO 2025: HNSCC swimmer plots (n=58)



CRB-701 biomarker populations: Observed efficacy

Nectin-4

Responses seen across wide range of IHC H-score expressions

HPV

Responses seen in HPV positive and negative patients

PD(L)-1

Responses in PD(L)-1 positive and negative patients

Case Study #1: Clinical improvement in participant with resistant disease

Prior therapies

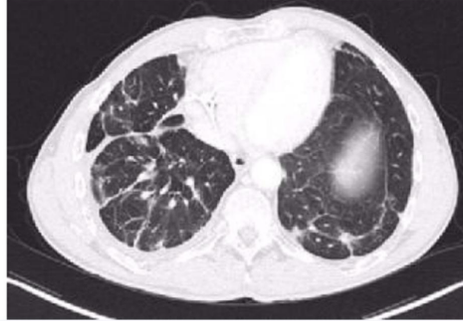
Carboplatin+docetaxel+5FU 3 weeks (PD)
then Cisplatin 4 weeks (PD) then
pembrolizumab 6 weeks (PD)
then experimental bispecific antibody (PD)

“ 61-year-old male patient with HNSCC PD-L1 <1 recently had 1 year tumor assessment images. He was previously suffering with significantly reduced performance status (ECOG 2) and on supplemental oxygen, now riding his bicycle, off oxygen and has gained 15 pounds with an ECOG of 0. ”

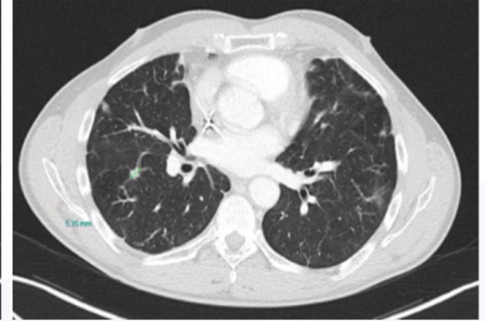
- USA Study Physician



Baseline tumor assessment 9/19/2024



6-week follow-up assessment 11/7/2024



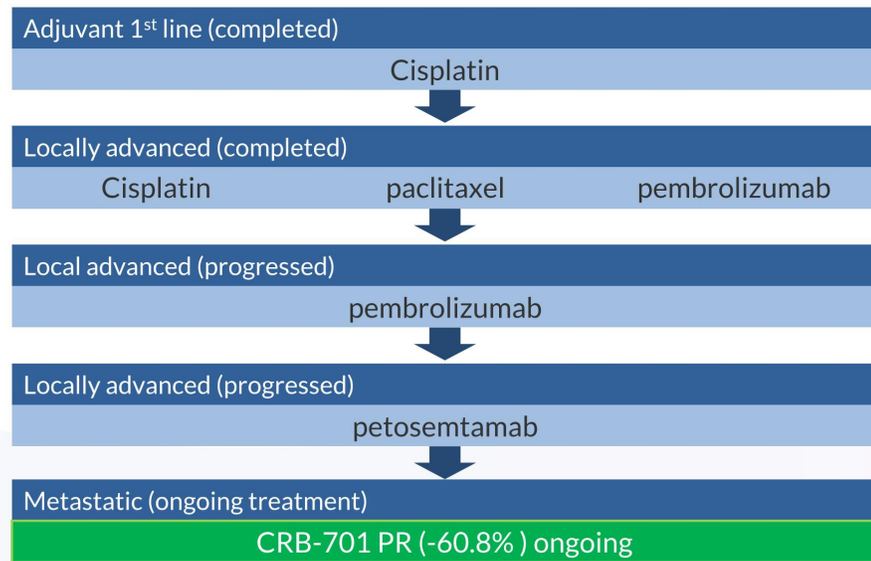
1-year follow up assessment 9/22/2025

As of 22 Sep 2025 – Participant is ongoing with a PR and tumor reduction of -73% with negative NavDx ctDNA. Remaining disease is PET negative/cold – being considered as a clinical (not formal) CR.

Case Study #2: Response seen in patient pre-treated with petosemtamab

Patient had a partial response (after previously showing stable disease while on petosemtamab)

Patient was heavily pre-treated with 4 lines of prior therapy



CRB-701 compared to petosemtamab or PADCEV® in 2L HNSCC

	Petosemtamab***	HNSCC PADCEV®**	CRB-701*
Dosing regimen	1500mg Q2W	1.25mg/kg on d1/8/15 of 28-day	3.6mg/kg Q3W
Target population	PD(L)-1 +ve only (HPV+/-)	PD(L)-1 agnostic (HPV+/-)	PD(L)-1 agnostic (HPV+/-)
Efficacy (ORR)	36%	23.9%	47.6%
TEAEs Grade 3 & greater	59%	34.8%	35.5%

Target patient populations for CRB-701 in HNSCC

1L

- Multiple MOAs being evaluated
- CRB-701 combo data with pembrolizumab → expected mid-2026

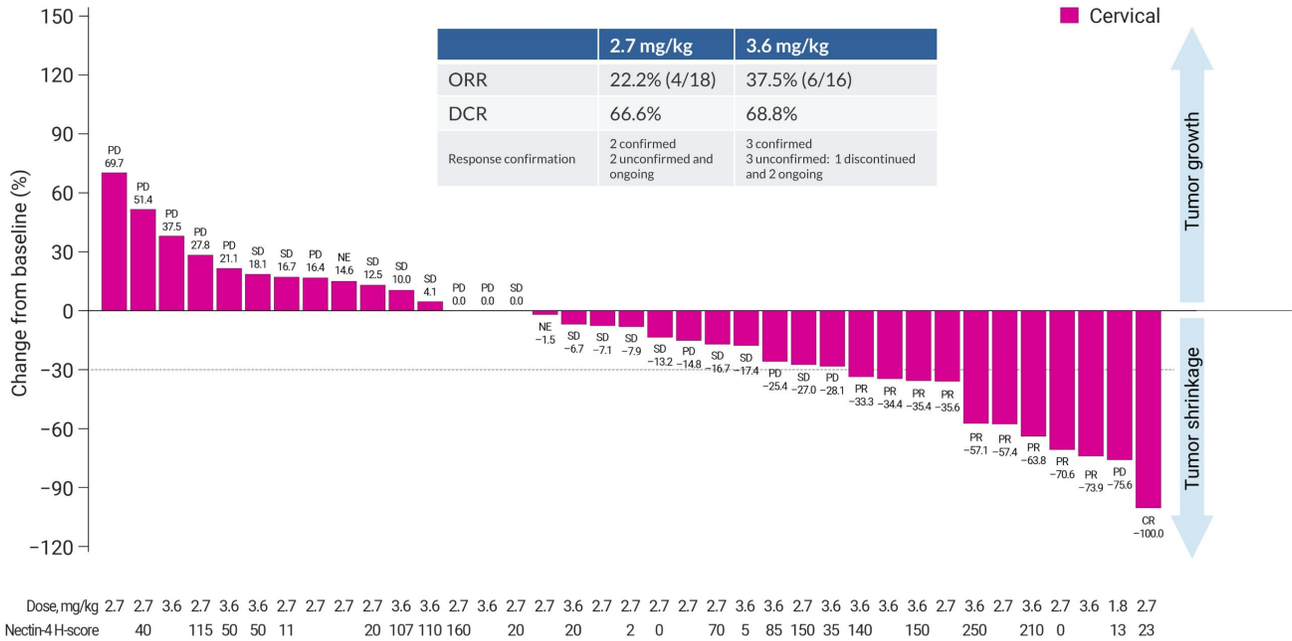
2L+

- 24,000* annual cases in USA
- No ADCs approved
- Orthogonal mechanism to EGFR
- Existing late line Tx ORR ~10-20%
- Petosemtamab ORR 36%

CRB-701 HNSCC: Next steps planned



ESMO 2025: Waterfall plot (n=37)



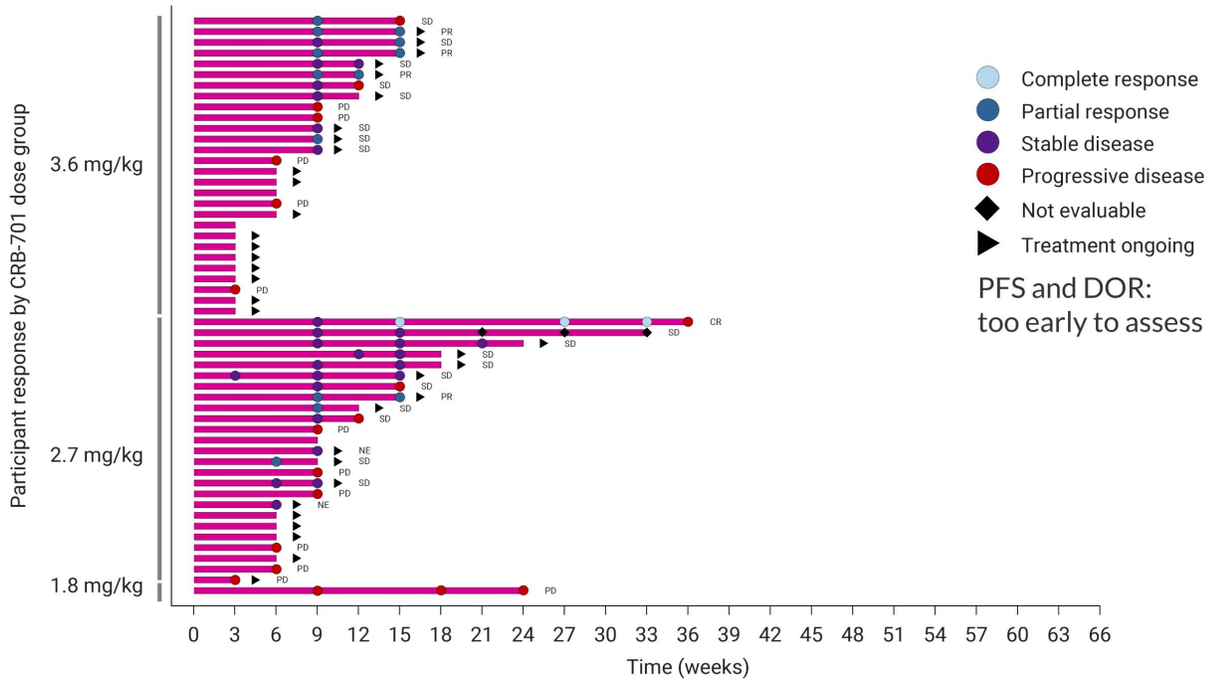
Source: ESMO 01 Sep 2025 Data cut
 Note: NE = Non-evaluable

ORR % = (CR+PR) / Response evaluable patients
 DCR % = (CR+PR+SD) / Response evaluable patients

3 patients excluded from ORR and DCR Calculations
 2 non-evaluable patients
 1 patient dosed at 1.8mg/kg



ESMO 2025: Swimmer plots (n=54)



ESMO 2025: CRB-701 compared to Tivdak®

	CRB-701	Tivdak®
Mechanism	Nectin-4 ADC with MMAE payload (DAR 2)	Tissue factor ADC with MMAE payload (DAR 4)
Target population	2L	2L
Median Age	54 (32-78)	51 (26-80)
ECOG (0, 1, 2, missing)	51.9%, 48.1%, 0%, 0%	61%, 39%, 0%, 0%
Prior lines of therapy median (range)	3 (1, 8)	1 line: 61% 2 lines: 38% Unknown: 1%
Dosing regimen	3.6 mg/kg Q3W	2 mg/kg Q3W
Efficacy (ORR)	37.5%	17.8%*
TEAEs Grade 3 & greater	35.5% (n=76)	46% (n=405)

Potential use of CRB-701 in cervical cancer

- Post-1L therapy represents unmet need with few effective modalities
- Tivdak® considered “a standard of care” in 2L with current annualized sales of \$314 million*
- Side effect profile + poor efficacy are limitations on Tivdak® commercial success
- FDA has granted CRB-701 Fast Track status in cervical cancer

1L**Keytruda® + chemo**

Efficacy (ORR ~68%**)

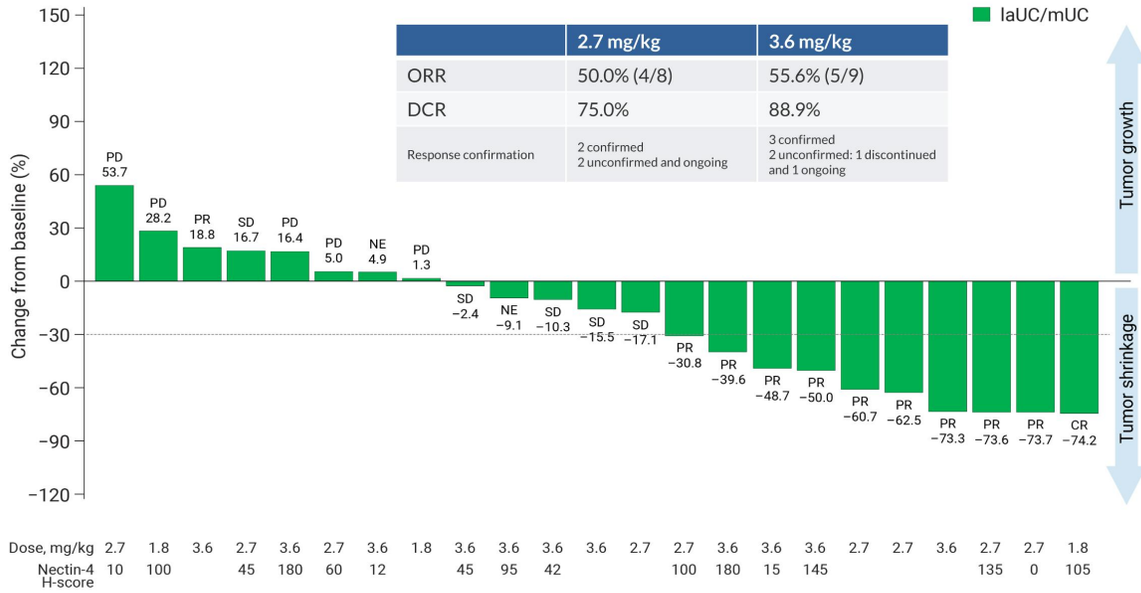
2L+**Tivdak®**

Modest efficacy (ORR 17.8%) and poor tolerability

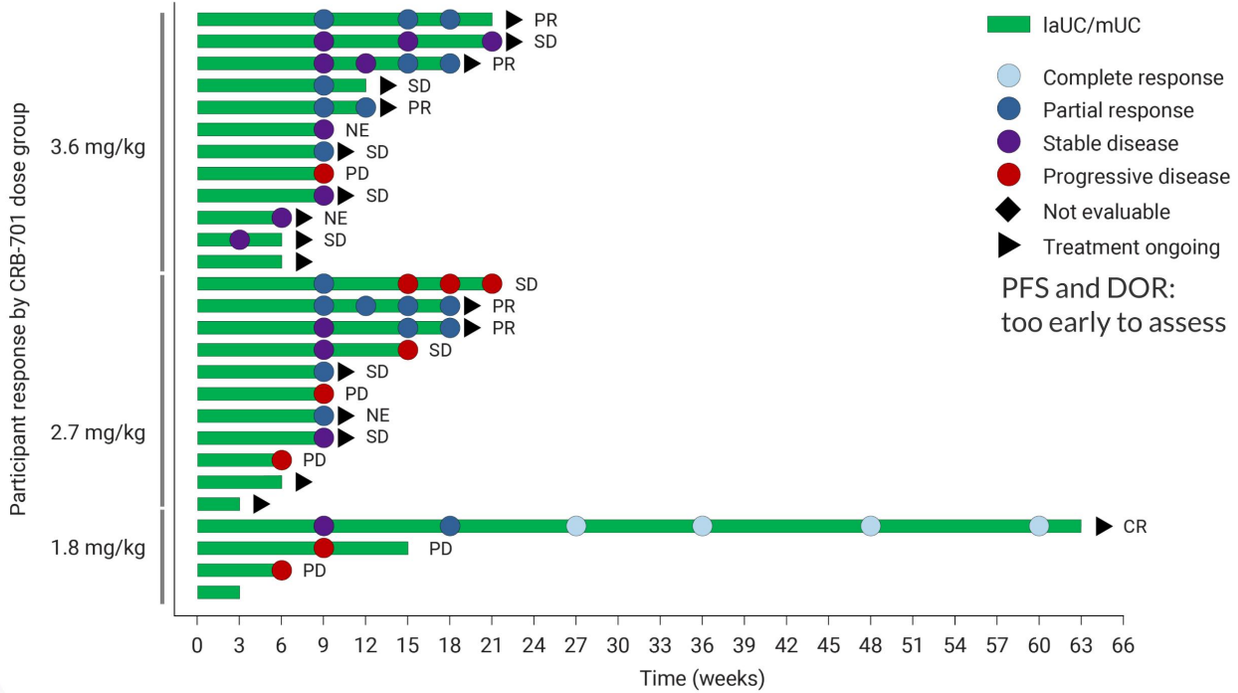
Source(s): *[Genmab Q2 YTD sales](#) of Tivdak® were \$78 million
*[Pfizer Q2 YTD sales](#) of Tivdak® were \$79 million

**[Keytruda prescription label](#)-Keynote 826 study

ESMO 2025: Waterfall plot (n=23)



ESMO 2025 Swimmer plots (n=27)



ESMO 2025: CRB-701 compared to PADCEV[®] monotherapy

	CRB-701*	PADCEV [®] **
Mechanism	Nectin-4 ADC with MMAE payload (DAR 2)	Nectin-4 ADC with MMAE payload (DAR ~3.8)
Dosing regimen	3.6 mg/kg Q3W	1.25 mg/kg on d1/8/15 of 28-day cycle
Target population	2 nd line	2 nd line
Efficacy-ORR	55.6%	44%
Pooled safety database	n=76	n=310 (1.25 mg/kg dose)
Grade 3 or greater AE rate	35.5%	58%
Peripheral neuropathy	6.6%	49%
Rash & skin reactions (broad terms)	29.3% (2.4% Grade 3 ^{***})	54% (7% Grade 3)
Discontinuation rates	7.9%	19.4%

Sources: *ESMO 01 Sep 2025 Data cut

**PADCEV[®] [data](#)

***All grade 3, no Grade 4/5: 1 x rash, 1 x decubitus ulcer, 1 x dermatitis bullous

Corbus not currently pursuing mUC as indication as a stand-alone company

- Decision based on current competitive landscape rather than data
- Keytruda® + PADCEV® dominate mUC 1L and PADCEV® dominates mUC 2L

1L

Keytruda® + PADCEV®

Efficacy:
(ORR 67.7%*)

2L+

PADCEV®**

Efficacy
(ORR 44%)

Sources: *Per [PADCEV® prescription label](#) EV-302 trial
**[PADCEV® data](#)

*

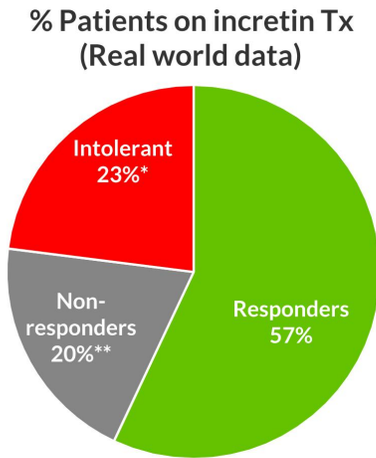
CRB-913

Daily oral small molecule targeting chronic obesity management

Data from Phase 1a SAD/MAD study

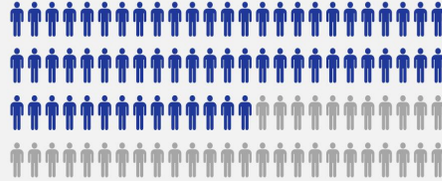


What are the emerging unmet needs in the obesity landscape?



64%

GLP-1 discontinuation @ 1 year
for obesity patients*



CRB-913's opportunity to reshape the obesity treatment paradigm



Alternatives to GLP-1 for resistant/intolerant/partial-responders



Lifelong weight maintenance using daily pill *post* weight loss



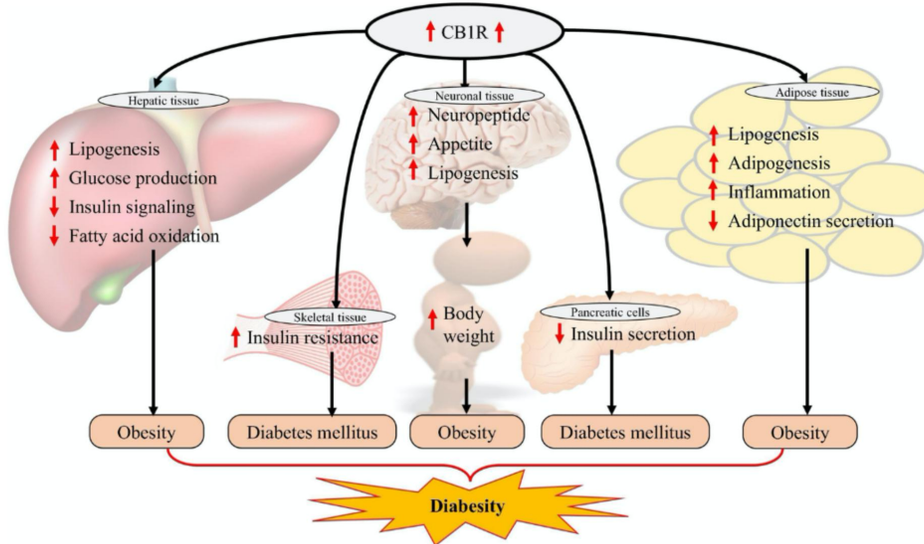
Avoiding sarcopenia

CB1 inverse agonism is only new non-incretin MOA that leads to weight loss

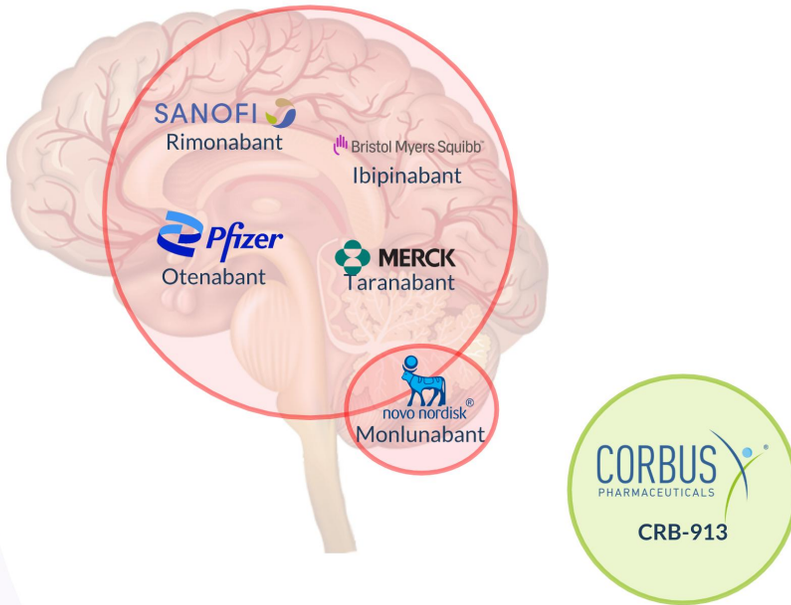
MOA	Company	Function	Monotherapy weight loss?
CB1 inverse agonism	Corbus, Novo	Appetite suppression, weight loss & muscle sparing	Yes
Activin	Lilly, Regeneron	Muscle sparing	No
Pan-agonist bitter taste receptor	Ardvark	Appetite suppression	No
INHBE siRNA	Wave, Arrowhead	Fat reduction + muscle buildup	No

CB1 is a well-understood receptor in metabolism

>9K papers in PubMed on CB1 and metabolism



CRB-913 is the first truly peripherally restricted inverse agonist



1/50th

Brain:plasma ratio
CRB-913 vs. Rimonabant*

1/15th

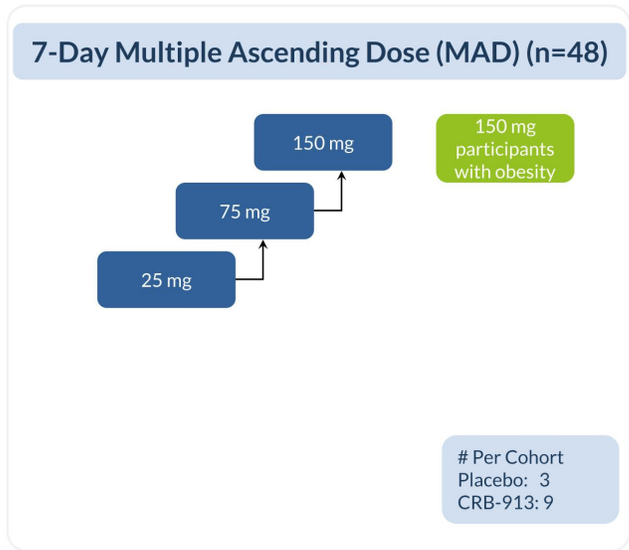
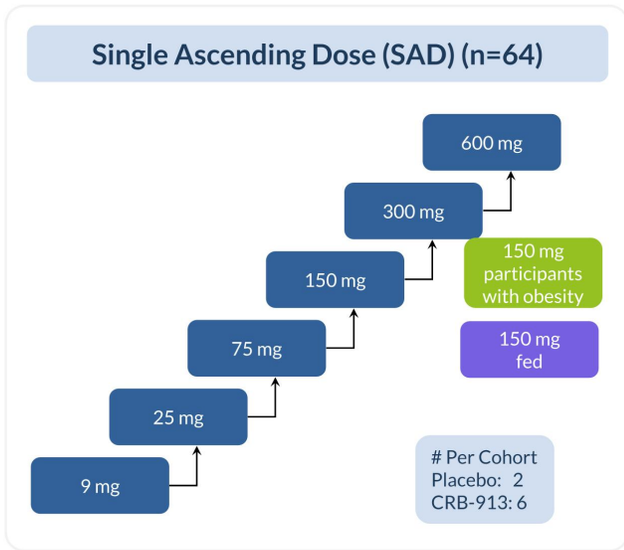
Brain level
CRB-913 vs. Monlunabant*

30%[↑]

Increase in peripheral levels in humans
vs. Monlunabant**



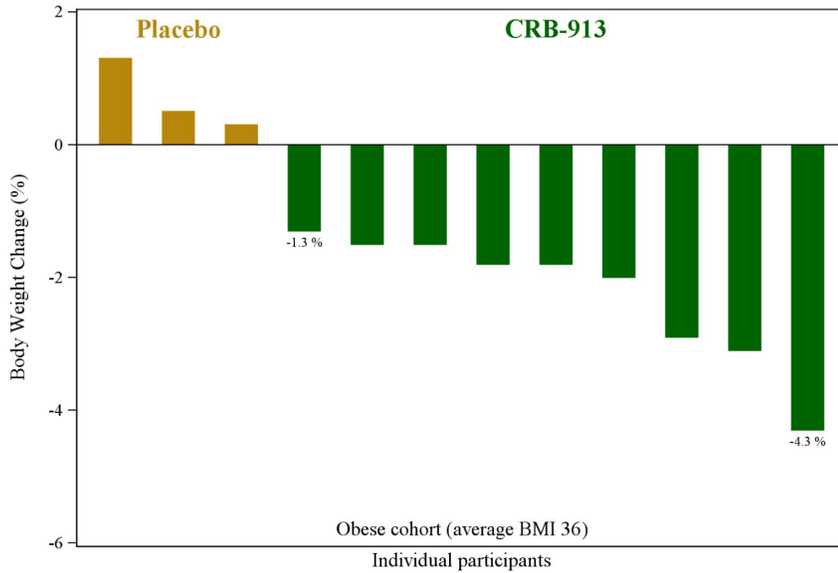
CRB-913 SAD/MAD study (Phase 1 unit in USA, total n=112)



Context:
Rimonabant efficacious dose: 20 mg QD
Monlunabant efficacious dose: 10 mg QD



Emerging weight loss with CRB-913 in subjects with obesity (150 mg MAD cohort)



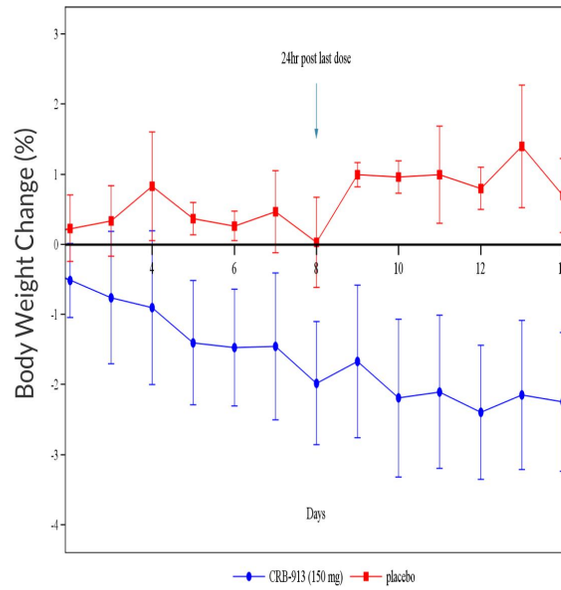
Participants reported reductions in food-related thoughts and cravings

2.9% average placebo-adjusted weight loss @ day 14

41 Note: Baseline is defined as the last available measurement taken prior to the first dose of study drug. Percent change in body weight is defined as body weight at Day 14 minus body weight at baseline divided by body weight at baseline multiplied by 100.

Weight loss with CRB-913 starts early and deepens

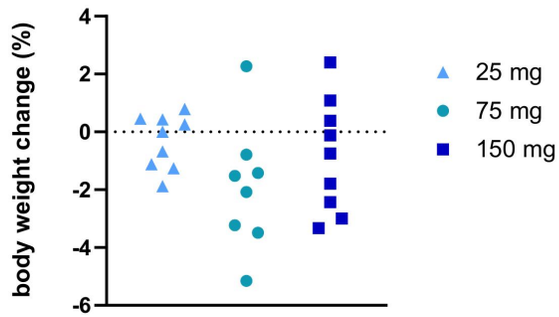
Obese cohort (average BMI 36) daily mean weight



42 Note: Baseline is defined as the last available measurement taken prior to the first dose of study drug. Percent change in body weight is defined as body weight at the given day minus body weight at baseline divided by body weight at baseline multiplied by 100.



Signals of weight loss in all-comer participants in MAD cohorts at lower doses



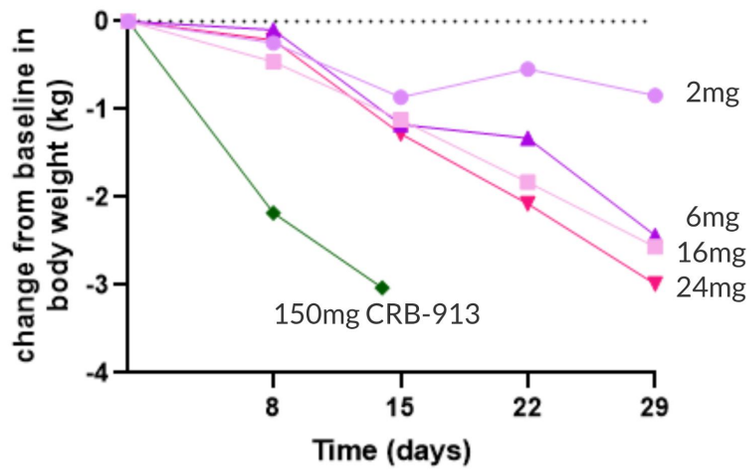
Important:
Weight loss for 75 mg QD
similar to 150 mg QD

Placebo-adjusted weight loss	▲ 0%	● 2.0%	■ 1.5%
BMI range	23.5 to 31.4	22.3 to 31.8	24.4 to 31.3

Average BMI of 28 → lower potential for weight loss

Emerging data CRB-913 vs. orfoglipron MAD: deeper and faster weight loss?

Placebo-adjusted weight loss cross-trial comparison for MAD studies



CRB-913 vs. orfoglipron MAD data: differentiated emerging safety

Adverse event	CRB-913*	Orfoglipron**
GI tolerability		
Nausea	None	12%-22%
Constipation	None	11%-23%
Vomiting	None	0%-18%
Neuropsych		
CSSRS	Negative	Negative
PHQ-9	Negative	Negative
GAD-7	Negative	Negative

* CRB-913 SAD/MAD data **Pratt et al 2023, Data is derived from cross-trial comparison.



Contextualizing weight loss in 2 weeks across oral MAD obesity clinical data sets

Drug	Company	placebo adjusted WL (%)	Type
CRB-913 (150 mg)		-2.9%	small molecule
Orforglipron (2 mg)		-1.4%	small molecule
Aleniglipron (5 mg)		-1.3%	small molecule
Elecglipton (50 mg)		0%	small molecule
Semaglutide (40 mg)		-0.7%	oral peptide
VK2735 (30 mg)		-1.8%	oral peptide

Potential clinical usage and supportive clinical or pre-clinical data (1 of 3)

1 CRB-913 monotherapy therapy for incretin insensitive/intolerant / high-risk patients



Insensitive patients: only hope is non-incretin MOA



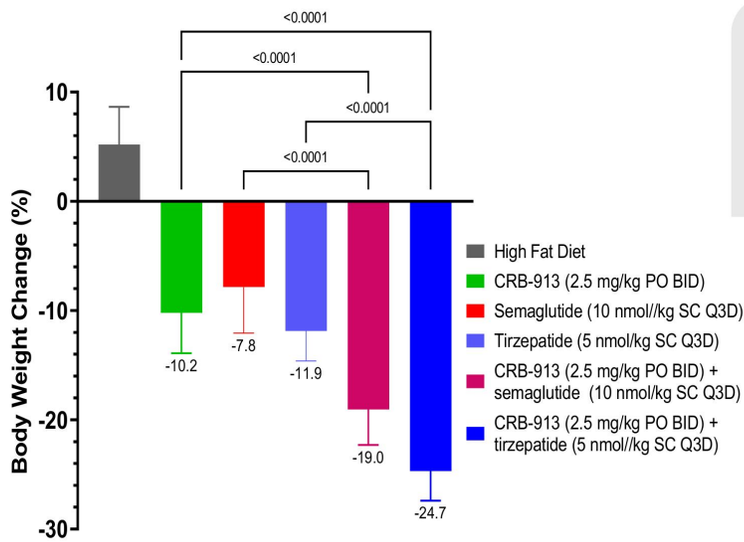
Intolerant patients : CRB-913 has markedly mild GI tox



High risk patients (sarcopenia): CB1 inverse agonism not associated with sarcopenia

Potential clinical usage and supportive clinical or pre-clinical data (2 of 3)

2 Combination with oral incretin agonists → potentially enhances efficacy OR improve tolerability



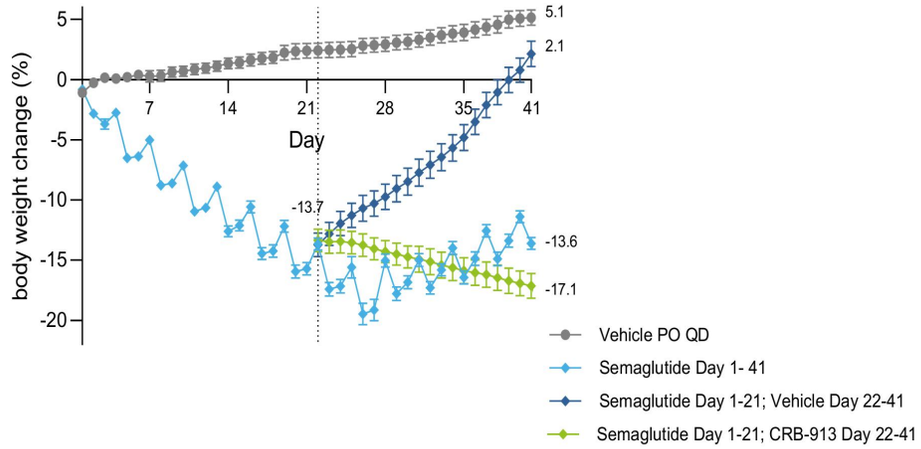
OBESITY SYMPOSIUM
Obesity Biology and Integrated Physiology

Obesity THE JOURNAL OF WILEY

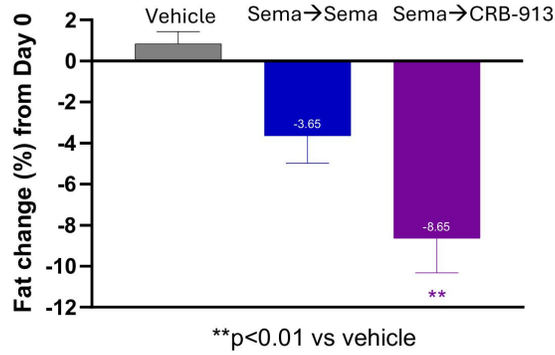
Novel cannabinoid receptor 1 inverse agonist CRB-913 enhances efficacy of tirzepatide, semaglutide, and liraglutide in the diet-induced obesity mouse model

Potential clinical usage and supportive clinical or pre-clinical data (3 of 3)

3 “Induction/maintenance” model: goal to potentially maintain weight loss post incretin analog therapy



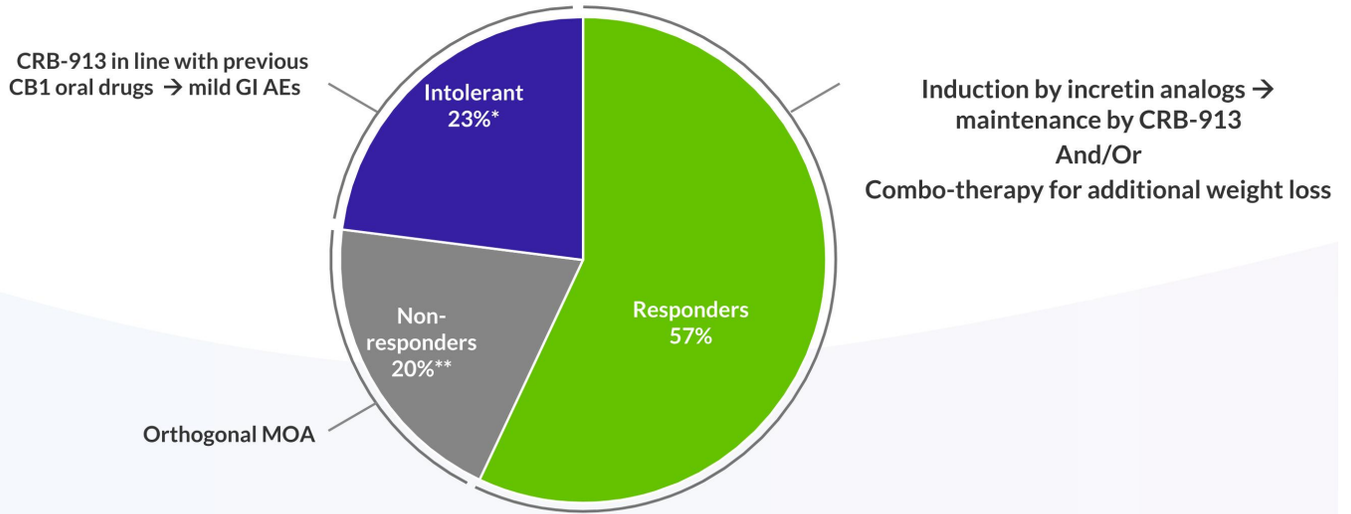
Weight loss from CRB-913 driven by further fat loss following semaglutide in DIO mouse model



At day 41 (end of study period)			
	Sema → Sema	Sema → CRB-913	Difference
Weight loss (%)	-13.6	-17.1	↑25%
Fat change from baseline	-3.65%	-8.65%	↑x2.3

What could the addressable market opportunity look like for CRB-913?

% Patients on incretin Tx (Real world data)

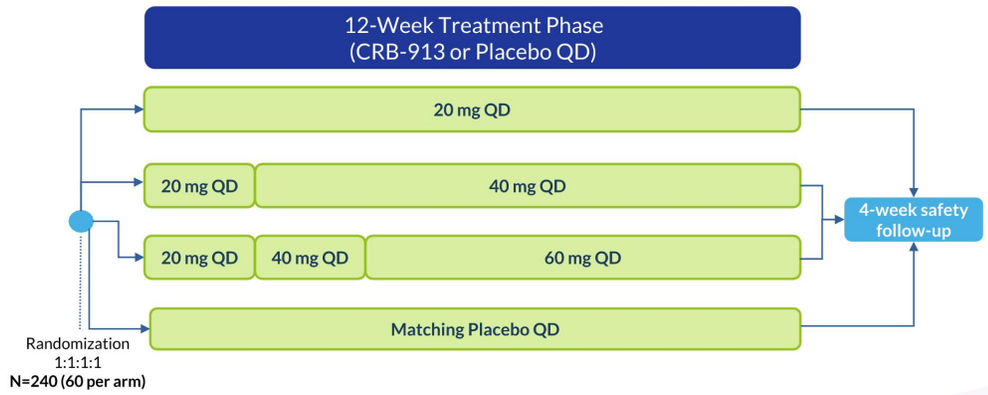


51 Source: *Cartwright et al 2025, **AP Nov 2024

Initiated: Phase 1b study



Completion summer 2026



	CRB-913 phase 1b (CANYON-1)	Monlunabant phase 2a
Subjects with obesity	240	240
Location	USA	Canada
Cohorts (all QD)	Placebo, 20, 40 and 60 mg	Placebo, 10, 20 and 50 mg
Titration	Yes	No
Exclude PHQ-9 > 4 at baseline	Yes	No

What did we learn from the CRB-913 SAD/MAD data?



True peripheral restriction → favorable safety + tolerability



CRB-913 elicits weight loss that starts early and deepens



Weight loss is *not* driven by GI AEs



Weight loss is associated with restriction to the periphery

CRB-601

Potential “best-in-class” $\alpha\beta 8$ mAb

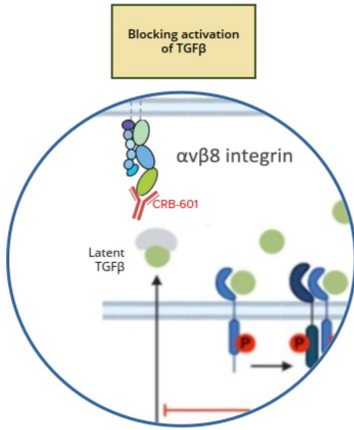
CRB-601 has the potential to enhance checkpoint inhibition



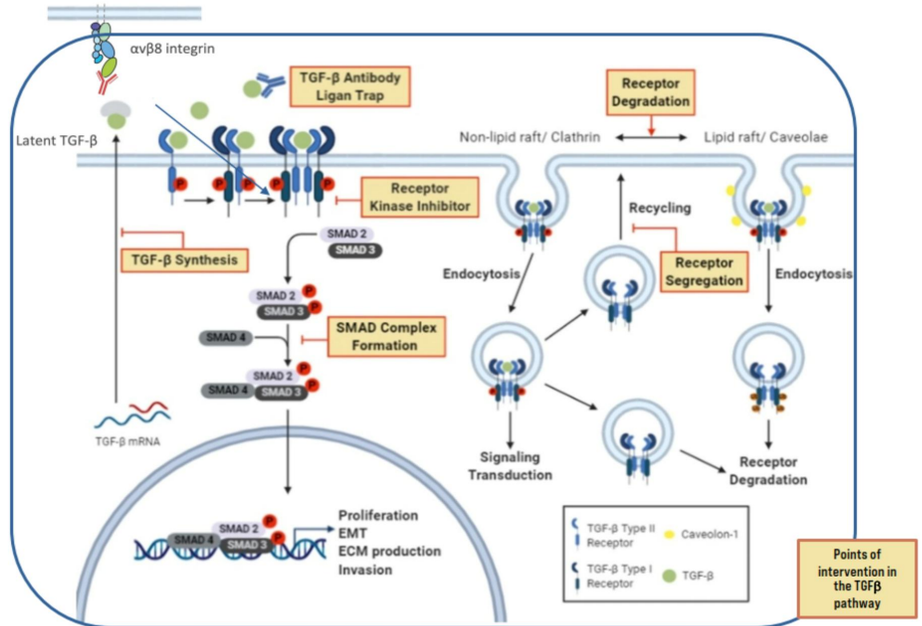
Targeting the integrin $\alpha\beta 8$ represents a novel approach to regulating TGF β

Novel point of therapeutic intervention

Blocking the $\alpha\beta 8$ activation of TGF β in the local tumor microenvironment



CRB-601 binds at the interface between latent TGF β and $\alpha\beta 8$



mAbs targeting TGFβ activation in the clinic

					
	CRB-601	PF-06940434	SRK-181	ABBV-151	RG6440
MOA	αvβ8	αvβ8	L-TGFβ	GARP (TGFβ1)	L-TGFβ
Clinical Stage	Phase 1	Phase 1/2 – study completed December 2024	Phase 1	Phase 2 HCC (read-out in 2025) Expanded Ph2 trials into muC & NSCLC	Phase 1
Indications	Solid Tumors	Solid Tumors	Solid Tumors	HCC	Solid Tumors
Type	Monoclonal Antibody	Monoclonal Antibody	Monoclonal Antibody	Monoclonal Antibody	Monoclonal Antibody
ROA	IV	IV	IV	IV	IV



Leadership
Upcoming Catalysts
Financials



Management team



Yuval Cohen, PhD
Chief Executive Officer,
Director

Corbus co-founder and Chief Executive Officer since 2014. Previously the President and co-founder of Celsus Therapeutics from 2005.



Sean Moran, CPA, MBA
Chief Financial Officer

Corbus co-founder and Chief Financial Officer since 2014. Prior senior financial management experience in emerging biotech and medical device companies.



Dominic Smethurst
Chief Medical Officer,
MA MRCP

Dr. Smethurst, MA MRCP, joined Corbus as our Chief Medical Officer in February 2024. He most recently served as CMO of Bicycle Therapeutics.



Ian Hodgson, PhD
Chief Operating Officer

Dr. Hodgson joined Corbus in 2022. Previously he held senior leadership positions in biotech and contract research organizations. Most recently served as V.P., Head of Clinical Services at TMC Pharma.



Christina Bertsch, M.A.
Head of Human Resources

Accomplished senior human resource executive providing strategic HR consulting services to both large and small businesses across a variety of industries.

Board of Directors



Rachelle Jacques **Chair of the Board**

More than 30-year professional career, experience in U.S. and global biopharmaceutical commercial leadership, including multiple high-profile product launches in rare diseases; Former CEO of Enzyvant Therapeutics (now Sumitomo Pharma) and Akari Therapeutics (NASDAQ: AKTX)



Anne Altmeyer, PhD, MBA, MPH **Director**

Greater than 25 years of experience advancing oncology R&D programs and leading impactful corporate development transactions; former CEO of TigeTx (acquired by Epsilogen Ltd)



Winston Kung, MBA **Director**

More than 20 years of senior financial, business development and investment banking experience; currently CFO of ArriVent. (NASDAQ: AVBP)



John K. Jenkins, MD **Director**

Distinguished 25-year career serving at the U.S. FDA, including 15 years of senior leadership in CDER and OND.



Yong (Ben) Ben, MD, MBA **Director**

25 years of oncology R&D experience across industry and academia. CMO of BridgeBio Oncology Therapeutics and former CMO of BeiGene.



Yuval Cohen, PhD **Chief Executive Officer, Director**

Corbus co-founder and Chief Executive Officer since 2014. Previously the President and co-founder of Celsus Therapeutics from 2005.

Upcoming anticipated corporate milestones

CRB-701	Regulatory update	Q1 2026
	Start HNSCC monotherapy Ph2/3 registrational study	Mid-2026
	Phase 1/2 monotherapy data	Mid-2026
	CRB-701+ pembrolizumab data	2 nd half 2026
CRB-913	Complete Ph1 SAD/MAD	Q4 2025 ✓
	Start Ph1B study	Q4 2025 ✓
	Complete Phase 1B	Summer 2026
CRB-601	Ph1 dose escalation	Q1 2026