

Propanc Biopharma Inc. OTC Pink: PPCB Nov 2023

www.propanc.com

Overview

Propanc Biopharma is a clinical stage biopharmaceutical company developing new cancer treatments for solid tumors. Propanc has developed a formulation of anti-cancer compounds designed to control or prevent tumors from recurring and spreading throughout the body by using proenzymes, which are inactive precursors of enzymes.

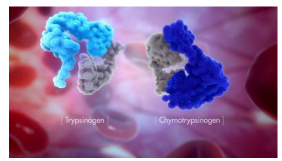
Propanc intends to target patients with limited therapeutic options for treatment of solid tumors, initially pancreatic, ovarian or colorectal tumors. Propanc achieved Orphan Drug Designation from the FDA for the treatment of pancreatic cancer. Propanc is also developing its lead product, PRP, to treat early stage cancer and pre-cancerous diseases and as a preventative measure for patients at risk of developing cancer, based on genetic screening.

Stock Data

Price (11-21-23)	\$0.0112
Market Cap	\$249.30
Avg. Daily Volume (30D)	446,899
Outstanding Shares	22,258,941
52 week High and Low	\$1.10/ \$0.0007

PRP is designed to eradicate Cancer Stem Cells

- Mixture of two proenzymes, trypsinogen & chymotrypsinogen from the bovine pancreas.
- PRP induces cell differentiation, converting cancerous cells into normal cells.
- Compassionate use patient data shows statistically significant results.
- PRP is a patented approach that:
 - Suppresses tumor metastasis and relapse.
 - ➤ Complements conventional anti-cancer therapies.
 - ➤ Is safe at specified dosages with minimal toxicity.
 - ➤ Is not cytotoxic (toxic to living cells)
- A synergistic ratio of 1:6 inhibits growth of most tumor cells.
- Examples include ovarian and colorectal cancers.
- Has also shown efficacy in kidney, breast, brain, prostate, lung, liver, uterine, and skin cancers.



Investment Highlights

Targeted Therapy for Metastatic Cancer:

No effective standard treatments exist for solid tumors. Tumor cells return to the normal pathways of a differentiated cell.

Multiple Mechanisms of Action

PRP is a novel technology with a large scale global potential that not only treats metastatic cancer but reduces the rate of recurrence.

Unique Intellectual Property

The Company is building a robust patent portfolio around its scientific understanding of the effects of pro-enzymes in cancer, new formulations, new routes of administration, and potential new targets.

Expansive Market Demand:

Approximately 80% of cancers are from solid tumors, and metastasis is the main cause of patient death. Proenzyme therapy targets and eradicates cancer stem cells not killed by radiation or chemotherapy.

Encouraging Patient Data

15 years of scientific research and clinical experience suggest PRP has the potential to convert cancerous cells back into normal cells.

International Strategic Partnerships

The Company maintains research partnerships with the University of Bath, University of Jaén, University of Granada and the Foundation for Biosanitary Research of Andalusia Orients.



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Compassionate Use Data

- 46 terminal patients (UK & AUS) administered two proenzymes plus amylase via suppository.
- Independent review concluded 19 patients significantly exceeded life expectancy.
- Response rate comparable to cytotoxic or immunologic approaches at Phase 1.
- No severe or even serious adverse effects.
- Most showed improved quality of life and relief of symptoms.
- Increased dose may result in better therapeutic efficacy.

Propanc Innovation & Intellectual Property

- Sixty five patents in force and pending covering several important discoveries regarding proenzymes and their anti-cancer effects:
 - Pharmaceutical composition for treating cancer
 - ➤ Proenzyme compositions
 - ➤ Cancer treatment (eradicating CSCs)
 - Composition of proenzymes for cancer treatment
- Lead patent approved in several countries including the US & EU.

PRP Development Timeline

	' 24			' 25			
	Q1	Q2	Q3	Q4	Q1	Q2	Q3
GMP Manufacture / Bioanalyticals							
Obtain Regulatory Approval for Phase IB							
Phase IB, F.I.H. study							

	Mr James Nathanielsz		Dr Julian Kenyon		Prof Klaus Kutz	
	Chief Executive Officer		Chief Scientific Officer		Chief Medical Officer	
•	Director and Chief Executive	•	Founded the company and	•	20 years of experience as	
	Officer since October 2007		appointed Director on February		independent consultant in	
•	20 years of experience in R&D,		12, 2008		clinical pharmacology and safety	
	manufacturing, and distribution	•	Medical Director of the Dove		in oncology for pharmaceutical	
	including 10 years in oncology		Clinic for Integrated Medicine,		companies and clinical research	
	and the development of		UK since 2000		organizations	
	chemotherapeutics	•	Bachelor of Medicine and	•	12 years of experience as Head of	
•	Bachelor of Applied Science,		Surgery and Doctor of		Pharmacology in 2 multinational	
	(Biochemistry/ Applied Chemistry)		Medicine, University of		pharmaceutical companies	
	and Master of Entrepreneurship &		Liverpool	•	Specialist for Internal Medicine,	
	Innovation, Swinburne University of	•	Primary Fellow of the Royal		Gastroenterology, and Clinical	
	Technology, Melbourne, Australia		College of Surgeons, Edinburgh		Pharmacology	
			for over 40 years	•	Professor of Medicine, University	
					of Bonn, Germany	

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