

Quantitative systems pharmacology modeling of biotherapeutics in oncology

Betts, A.M.

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Section VI. Appendices

Curriculum vitae

Alison Betts (1972, Edinburgh, Scotland) began her scientific career in 1990 at the University of St. Andrews in Scotland. She graduated in 1994 with a first-class honors degree in Biochemistry (BSc.) and was recipient of the class prize in Biochemistry. This course included an internship in an immunology research group at St. Andrews. In 1994, Alison moved from Scotland to the south of England to start her career at Pfizer in the department of Drug Metabolism. Here she gained extensive hands on experience of drug metabolism and pharmacokinetics and supported many small molecule drug discovery projects across diverse therapeutic areas.

From 2002-2004, Alison completed a secondment in the clinical pharmacometrics group at Pfizer. Here she learned the art and science of modeling, providing PK/PD support to the Pain research unit. In 2005, Alison joined the newly formed preclinical PK/PD group under the leadership of Dr. Piet van der Graaf. In this role she supported the Genitourinary and Obesity research units, using PK/PD modeling in the design and interpretation of experiments for small molecule drugs and performing PK and PK/PD predictions to human.

In September 2007, Alison moved to the USA with her family, to take up a new opportunity working in the Translational Modeling and Simulation (TMS) group in the department of Pharmacokinetics, Dynamics, and Metabolism (PDM) in Groton, CT. In this role, Alison had a diverse and extensive modeling career, supporting small and large molecule mathematical modeling across many research areas. Highlights included her role as TMS team leader supporting the antibacterial research group. In this role she was responsible for the successful execution of PK/PD modeling and translational strategy across the antibacterial portfolio including projects from exploratory stage through to Phase 1 clinical development. Here, Alison implemented mechanism-based PK/PD, PBPK and QSP models to predict efficacy and emergence of resistance and used these to develop novel strategies for clinical dose predictions of antibacterial drugs. In this work, Alison's team was awarded a Pfizer Leadership Team Excellence Award 2009. A particular focus was the establishment of *in vitro*: *in vivo* correlations so that *in vitro* models could be used to replace *in vivo* experiments. This work resulted in a reduction in *in vivo* animal spend by 40% and was awarded Pfizer Animal Care and Welfare Board 'Reduce, Re-use and Refine' award in 2010.

In 2010, Alison assumed responsibility as TMS team leader supporting the biotherapeutics division of the Oncology research unit at Pfizer. Alison's team supported a diversity of novel biotherapeutic modalities including proteins, peptides, antibodies, antibody drug conjugates (ADCs), bispecific antibodies, T cell engagers, cytokine conjugates and nanoparticles. Highlights of this work included providing modeling and simulation support to progress two ADCs to the market (Mylotarg® and Besponsa®) and many others to clinical studies. In this time, Alison was leader of the ADC working group at Pfizer, defining the quantitative analytical and predictive strategy for ADCs at Pfizer from preclinical stage through to clinical studies and she completed a

successful external partnership with Prof. Dane Wittrup's lab at the Koch Institute for Integrative Cancer Research, MIT to model intracellular trafficking of ADCs.

During this time, Alison became leader of the preclinical modeling and simulation discipline for the biotherapeutics division of PDM, including TMS teams in Cambridge MA, Groton CT and La Jolla CA. This included overseeing TMS support to Immunology & Inflammation, Oncology, Rare Diseases, Cardiovascular metabolic diseases, Neuroscience, and the Center for Therapeutic Innovation. Here she was responsible for leading scientific vision, defining strategy and for day-to-day process management of the group, including technical, infrastructure, resourcing (internal and outsourcing) and budgetary requirements. In this time, Alison's role and team transitioned from Groton, CT to Cambridge, MA.

In 2017, during her time at Pfizer, Alison started her PhD research on quantitative systems pharmacology modeling of biotherapeutic drugs in oncology at the Leiden Academic Centre for Drug Research (LACDR), under the supervision of Prof. dr. Piet H. van der Graaf. This gave Alison the opportunity to further explore her passions for using mechanistic modeling to enable quantitative decision making in oncology drug discovery, for preclinical to clinical translation, to optimize clinical dosing regimens and address precision medicine questions.

After 25 years at Pfizer, Alison started a new career path with Applied Biomath in 2019. Alison is Senior Director of Scientific Collaborations and Fellow of Modeling and Simulation at ABM, where she collaborates with partners to introduce mechanistic system pharmacology modeling approaches at preclinical and clinical stages to de-risk drug programs within the pharmaceutical and biotechnology industry. She maintains a hands-on approach as project leader for several modeling collaborations across therapeutic modalities. She is also recipient and Principal Investigator of an NIH SBIR Grant to build a platform QSP model for ADCs (ADC Workbench) capable of predicting efficacy and toxicity with the aim of reducing TI.

In her diverse and extensive modeling and simulation career, Alison has published 31 manuscripts and has given 40 invited presentations at conferences.

List of Publications

- 1. Spinosa, P., Musial-Siwek, M., Presler, M., Betts, A., Rosentrater, E., Villali, J., Wille, L., Zhao, Y., McCaughtry, T., Subramanian, K., & Liu, H. (2021). Quantitative modeling predicts competitive advantages of a next generation anti-NKG2A monoclonal antibody over monalizumab for the treatment of cancer. *CPT: pharmacometrics & systems pharmacology*, 10(3), 220–229. https://doi.org/10.1002/psp4.12592
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Affiliations of Authors

Lindsay B. Avery

Department of Biomedicine Design, Pfizer Inc., 1 Burtt Road, Andover, MA

Frank Barletta

Oncology Research & Development, Pfizer Inc, 401 N Middletown Rd, Pearl

River, NY 10965

Jan Berkhout LAP&P, Archimedesweg 31, 2333 CM Leiden, The Netherlands

Joseph Boni Department of Clinical Pharmacology, Pfizer Global Research and

Development, Collegeville, PA

Subramanyam Chakrapani Department of Worldwide Medicinal Chemistry, Pfizer Global Research and

Development, Groton, CT 06340

Xiaoying Chen Department of Clinical Pharmacology, 10555 Science Center Dr, San Diego

CA 92121

Tracey Clark Established Med Business, Pfizer Inc., Eastern Point Rd, Groton CT 06340
Piet H. van der Graaf Division of Systems Biomedicine and Pharmacology, Leiden Academic

Centre for Drug Research, 2300 RA Leiden, The Netherlands

Edmund I. Graziani Apertor Labs Inc., 828 Contra Costa Ave., Berkeley, CA 94707

Nahor Haddish-Berhane Clinical Pharmacology and Pharmacometrics, Quantitative Sciences,

Janssen Pharmaceuticals, Spring House, PA 19002

Steve Hansel Translational Research Group, Department of Pharmacokinetics Dynamics

and Metabolism, Pfizer Global Research and Development, Groton, CT

06340

Andrea Hooper Oncology Research & Development, Pfizer Inc, 401 N Middletown Rd, Pearl

River, NY 10965

Paul Jasper RES Group, Inc., 75 Second Avenue, Needham, MA 02494

Theodore R. Johnson Department of Pharmacokinetics Dynamics and Metabolism, Pfizer Global

Research and Development, La Jolla, CA

Hannah Jones Department of Biomedicine Design, Pfizer Inc., 610 Main Street, Cambridge,

MA

Cris Kamperschroer Department of Immunotoxicology, Pfizer Inc, 558 Eastern Point Road,

Groton CT 06340

Anne Keunecke LAP&P, Archimedesweg 31, 2333 CM Leiden, The Netherlands

Lindsay E. King Department of Biomedicine Design, Pfizer Inc., 1 Burtt Road, Andover, MA
Frank Loganzo Oncology Research Unit, Pfizer Global Research and Development, Pearl

River, NY 10965

Judy Lucas Oncology Research & Development, Pfizer Inc, 401 N Middletown Rd, Pearl

River, NY 10965

Dangshe Ma Oncology Research & Development, Pfizer Inc, 401 N Middletown Rd, Pearl

River, NY 10965

Sylvia Musto Oncology Research & Development, Pfizer Inc, 401 N Middletown Rd, Pearl

River, NY 10965

Adam Root Department of Biomedicine Design, Pfizer Inc, 610 Main Street, Cambridge,

MA 02139

Edward Rosfjord Oncology Research & Development, Pfizer Inc, 401 N Middletown Rd, Pearl

River, NY 10965

Dhaval K. Shah Department of Pharmaceutical Sciences, 455 Kapoor Hall, University at

Buffalo, The State University of New York, Buffalo, New York 14214-8033

Boris Shor Immune Pharmaceuticals Inc., 430 East 29th Street, Suite 940, New York,

NY 10016

Tamara J. van Steeg LAP&P, Archimedesweg 31, 2333 CM Leiden, The Netherlands

Matthew Sung Oncology Research & Development, Pfizer Inc, 401 N Middletown Rd,

Pearl River, NY 10965

John Tolsma RES Group, Inc., 75 Second Avenue, Needham, MA 02494

Hallie S. Wald Oncology Research & Development, Pfizer Inc, 401 N Middletown Rd,

Pearl River, NY 10965

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Finally, this thesis is dedicated to and inspired by my mother, Susan Harper, who died of metastatic breast cancer at the age of just 50. In her abbreviated time on this earth, she achieved a lot and has been a continuous source of inspiration. Working in cancer research has been an honor and a privilege, and this thesis is dedicated to her memory.

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