

UCB PharmD Fellowship Program Program Guide and Application Information 2025–2027





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Message from Executive Sponsors

At UCB, we believe that everyone deserves to live the best life that they can. That's why – as a global biopharmaceutical leader - we're focused on creating valuable solutions that make improvements to the lives of people living with neurological and autoimmune conditions now and into the future. We are Inspired by Patients. Driven by Science. These are not only words but are the cornerstone of our patient value culture at UCB. With a diverse portfolio of marketed products and a deep pipeline of new assets in discovery and early clinical stages, UCB measures success by the value we can deliver with our solutions.

As we continue to build on previous successes, we are committed to scientific innovation and organizational agility to keep pace with the evolving healthcare landscape. To succeed in this commitment, talent is key. We focus on developing talent with the competencies, skills, and capabilities needed to successfully deliver patient value in this complex environment.

The UCB PharmD Fellowship Program is designed to provide PharmD graduates with the opportunity to learn and experience all aspects of a specific functional area under the mentorship of experienced preceptors. As a mid-size pharmaceutical company, Fellows will have significant opportunity to interact with senior leaders at UCB, thereby enhancing their learning experience.

If you have the desire to work in a biopharmaceutical company with a focus on patient value, innovation and agility, and commitment to staff development, we encourage you to apply to the UCB Fellowship Program.

About UCB

UCB, founded in 1928 by Emmanuel Janssen, is a global biopharmaceutical company committed to developing innovative solutions to address significant unmet needs for people living with severe, chronic diseases. Our people, with their diversity, unique strengths, and talents, enable us to fulfill our commitment. With a team of approximately 9,000 employees and operations in nearly 40 countries, UCB is investing more than a quarter of its revenue in cutting-edge scientific research to meet unmet patient needs. Global headquarters are in Brussels, Belgium, with U.S. headquarters in Atlanta, Georgia. Additional U.S. UCB sites are located across California, Massachusetts, North Carolina, Washington, and Washington, D.C.

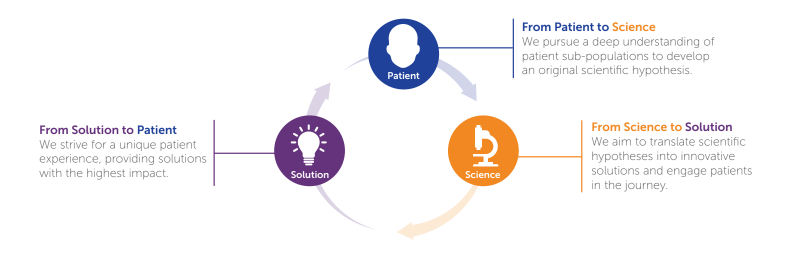
By putting **patients at the heart of everything we do,** we enable people to **live their best lives,** delivering impactful solutions **patients value.**



UCB Pipeline

At UCB, we want to help people live their best lives, whatever that means for them. We're focused on severe, chronic neurological and immunological conditions regardless of population size. And we're blazing a path integrating new technologies, like machine learning and data analytics, into how we work today to unlock a healthier tomorrow.

Our approach to innovation keeps patients at the center. We use patient insights to inform our science to build solutions to deliver to patients. Innovation is ongoing as we continue to search for solutions to meet the unmet needs of patients.



UCB's innovation delivering industry-leading pipeline

UCB is connecting science in new ways to illuminate the biological pathways involved in severe diseases. Our researchers are developing a range of novel chemical entities (NCEs) and novel biological entities (NBEs) to improve people's lives.

	PHASE 1	PHASE 2	PHASE 3	TOPLINE RESULTS
rozanolixizumab (FcRn inhibitor)				
MOG-antibody disease				H2 2026
Severe fibromyalgia syndrome		Ph-2a		H2 2024
fenfluramine (5-HT agonist)				
CDKL5 deficiency disorder				H2 2024
doxecitine and doxribtimine (nucleoside therapy)				
TK2 deficiency disorder				Submissions to begin end 2024
dapirolizumab pegol (anti-CD40L antibody)				
Systemic lupus erythematosus*				Mid-2024
STACCATO® alprazolam (benzodiazepine)			_	
Stereotypical prolonged seizures				H1 2026
bepranemab (anti-tau antibody)				
Alzheimer's disease**		Ph-2a		H2 2024
minzasolmin (a-syn-misfolding inhibitor)				
Parkinson's disease***		Ph-2a		H2 2024
UCB0022 (D1 receptor positive allosteric modulators)				
Parkinson's disease		Ph-2a		H1 2025
UCB9741				
Atopic dermatitis		Ph-2a		H2 2024
UCB1381			_	
Atopic dermatitis		Ph-2a		H2 2024

Inspired by patients. UCB - HY results 2024, July 2024. *In partnership with Biogen; 1st phase 3 study; **In partnership with Roche / Genentech; ***In partnership with Novartis; 5-HT = 5-hydroxytrytamin or serotonin; o-syn = alpha-synuclein; CD40L = CD40 ligand; CDKL5 = cyclin-dependent kinase-like 5; H = half-year; IL = interleukin; FcRn = Neonatal Fragment Crystallizable Receptor; MOG = Myelin Oligodendrocyte Glycoprotein; TK2 = Thymidine Kinase 2; projects not currently approved by any regulatory authority

UCB PharmD Fellowship Program

The UCB PharmD Fellowship Program, a collaboration with the Industry Pharmacists Organization (IPhO), is a 2-year program in the following functional areas:

Global Regulatory Affairs (GRA) Medical Safety and Pharmacovigilance (MS&PV) Medical Affairs – Immunology (MA) Global Clinical Sciences and Operations (GCSO) Global Medical Affairs (GMA)

The UCB Fellowships will be located at one of the following UCB campuses: the Atlanta campus in Smyrna, GA or the Research Triangle Park (RTP) campus in Morrisville, North Carolina. The GRA, MA and GMA Fellowships are stationed at the Atlanta campus. The GCSO Fellowship is located at the RTP campus. The MS&PV Fellow will have the option to choose between the Atlanta or the RTP campus.

What's unique about the UCB-IPhO Fellowship?

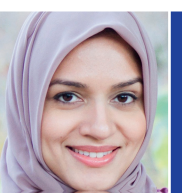
The UCB Fellowship Program offers a unique opportunity to work in an environment that is patient focused, creative, flexible, and agile, with an exciting and promising pipeline.

The support of Fellowship leadership, preceptors, and mentors, coupled with the unique combination of rotations and hands-on experiences, will help to ensure the success of the Fellows, developing them to become best-in-class industry professionals ready for a career in a variety of settings. Following two years, the Fellow will have the experience to move into a strategic/ operational (manager/ senior manager) role within the pharmaceutical industry, contract research organizations (CROs), or the FDA.

In addition, this Fellowship is offered in collaboration with IPhO. Through IPhO, the Fellow can gain exposure to broader networking and leadership opportunities for pharmacists in industry.

Benefits of the IPhO partnership include:

- Organizational Leadership: Fellows will be required to be members of the IPhO National Fellows Council (NFC) and will be given priority in holding leadership positions to develop and practice cross-functional leadership skills in the following committees: Fellows Development, Student Development, & Professional Programming.
- **Professional Development:** As a part of the IPhO NFC, Fellows will have access to fellow-targeted career development programming, such as webinars and live events.
- Publication Opportunities: Fellows can conduct research and/or publish a poster/paper/article in conjunction with an IPhO leadership team member
- Networking Opportunities: As a part of the IPhO NFC, Fellows will have the opportunity to network, both in-person and virtually, with Fellows across the country in various functional areas from both IPhO and non-IPhO Fellowship programs.
- **Teaching Experience:** Fellows will have an opportunity to be an instructor for IPhO Institute for Pharmaceutical Industry Learning (webinars), as well as provide guidance to hundreds of student pharmacists at 100 IPhO chapters.
- Mentorship: Fellows will receive mentorship from IPhO leadership.



"The UCB GRA Fellowship has been built to provide Fellows with a unique global regulatory experience – one that establishes a strong foundation in regulatory knowledge coupled with the autonomy to tailor the program to the Fellows' interests. With the support and mentorship of executive leadership and seasoned regulatory professionals across UCB and IPhO, the program positions Fellows on the path to a successful career in global regulatory affairs."

- Iram Hasan, Regulatory Science Lead, UCB GRA Fellowship Program Director

Recruiting 1 Fellow

The mission of Global Regulatory Affairs at UCB is to create innovative regulatory pathways and partnerships that expedite and maintain patient access to novel healthcare solutions. The GRA Fellowship provides Fellows with the depth and breadth of experience with all aspects of Regulatory Affairs to enable them to fulfill that mission and successfully position them for a career as a uniquely well-rounded Regulatory professional.

During the rotations within the sub-functions of Regulatory Affairs, the Fellows are assigned to work with the Regulatory Science Lead for one or more compounds, including pipeline and marketed products, ensuring that the chosen projects offer the greatest learning opportunity and exposure to FDA and other global regulatory health authorities. Additionally, the longitudinal exposure to Regulatory Operations throughout the two-year Fellowship provides the Fellows with a wholistic view of submission and project management. Lastly, the Fellows have an opportunity for a three-month elective in a functional area of their choice, outside of Regulatory Affairs, to allow them to gain additional insights from the outside in.

Rotation	Timeframe
Introduction to Regulatory Operations	2 week overlap with RTS
Regulatory Therapeutic Sciences (RTS)	8 months
Advertising-promotion & Labeling	5.5 months
Chemistry Manufacturing and Controls (CMC) & Devices	4.5 months
Regulatory Operations	1 month
Elective Rotation (outside GRA)	3 months
Final Core Rotation (within GRA)	2 months

Essential Functions & Responsibilities

- Support regulatory scientists/global regulatory leads in preparation and delivery of regulatory submissions, in collaboration with other support functions in GRA
- Support CMC associates to develop CMC-specific regulatory strategy and learn how to define content for CMC submissions
- Support advertising and promotion/labeling associates to understand regulatory requirements related to advertising and promotion as well as pharmaceutical company policies to ensure compliance with the regulations
- Acquire in-depth knowledge of fundamentals of regulatory affairs, regulatory intelligence, and development of regulatory strategy
- Provide regulatory operational support for pipeline and/or marketed product(s)
- Deliver project assignments supporting the business
- Develop proficiency in use of GRA systems



"UCB's Global Regulatory Affairs Fellowship stands out distinctly in post-doctoral training, offering unique opportunities through diverse rotations. These rotations empower fellows to lead projects while learning from world-class mentors and preceptors. The team members are enthusiastic about providing invaluable guidance and knowledge, nurturing each fellow's growth and development. UCB's culture exemplifies a commitment to patient-centric values, giving Fellows the ability to contribute meaningfully to patients' lives globally."

- Izzabella Christian, GRA 1st Year Fellow

As evident by the inclusive experience and the various rotations, UCB's Global Regulatory Affairs Fellowship is highly distinctive for post-doctoral training in this field. The program offers ample opportunity for leadership in meaningful projects, mentorship, and training with unique aspects of regulatory affairs such as devices and products for rare diseases. The diverse, welcoming, and truly patient-centered environment at UCB makes it an excellent place to learn and train to be an adaptable, impactful industry pharmacist."



Medical Safety and Pharmacovigilance Fellowship

Recruiting 1 Fellow

The UCB Medical Safety and Pharmacovigilance team provides end to end delivery and management of product benefit risk, safety profile, signal and risk management and strategic partnering as well as maintaining compliant safety reporting. These activities, in parallel with data transparency activities, build trust and enable new solutions to be delivered to patients and ensure the ongoing availability of our marketed products. Medical Safety and Pharmacovigilance function works transversally with multiple UCB stakeholders to deliver the above, both at a product level through leadership of the benefit risk team and more generally. The Medical Safety and Pharmacovigilance function work in both the scientific and operational disciplines that are required to ensure safety throughout a product's lifecycle.

Rotation	Timeframe
Product Pharmacovigilance & Device Safety	8 months
Safety Risk Management	8 months
International Pharmacovigilance (UCB affiliates)	3 months
Elective Rotation (outside MS&PV)	3 months
Final Core Rotation (within MS&PV)	2 months

Essential Functions & Responsibilities

- Actively contribute to safety aspects of designated products through project management, including global case processing, safety systems, signal detection, signal evaluation, benefit-risk assessments, case analysis, aggregate report creation, safety risk management contribution, and global health authority inquiry response, etc.
- Engage in reviewing UCB deliverables across multiple products and lifecycle stages, based on team needs ensuring alignment with safety management goals and standards.
- Collaborate with line managers, scientists, physicians, or equivalent on assigned tasks with Patient Value Solutions (PVSs), Established Brand Units (EBUs), clinical trials, submissions, etc.
- Support the IQF Governance Meeting for ICSR Quality Forum, upholding standards for Individual Case Safety Reports (ICSRs).
- Collaborate with the Medical Safety and Pharmacovigilance Leadership Team, actively sharing insights, ideas, and strategies for global patient safety enhancement.
- Lead or contribute to strategic initiatives to advance patient safety, leveraging expertise and UCB's vision.
- Gain an understanding of medical device safety and reporting, expanding skill set and comprehension.



"The UCB Medical Safety and Pharmacovigilance Fellowship Program offers fellows an outstanding opportunity to gain comprehensive exposure to the fundamentals and develop the skills needed to become a top-tier safety professional. The program ensures great visibility with the leadership team and a robust support system to ensure each fellow's success. Additionally, the program gives fellows opportunities to lead and contribute to significant projects, further enhancing their technical expertise and leadership skills. This enriching experience has been instrumental in shaping my career and professional growth. I am confident that UCB is preparing me to excel in the field."

- Maria Reji, MS&PV 1st Year Fellow

"The Medical Safety and Pharmacovigilance Fellowship program here at UCB will provide me with the skillset and fundamentals to be a well-versed pharmacovigilance scientist as I rotate through different areas of pharmacovigilance. This program is incredibly unique giving fellows' exposure to medical device safety and surveillance along with international pharmacovigilance. UCB is the epitome of a patient-centric company; they are committed to the well-being of patients which creates a dynamic atmosphere to learn and grow. Fellows have the opportunity to take on and lead projects and also gain mentorship from experienced professionals. In addition, the IPhO component allows for fellows to gain professional development opportunities."



- Collins Kofi Asamoah, MS&PV 2nd Year Fellow

Medical Affairs – Immunology Fellowship

Recruiting 2 Fellows

The Medical Affairs – Immunology Fellowship focuses on opportunities to learn, experience and lead various activities involved within the dynamic functions within a Medical Affairs organization. Fellows will utilize their first year to learn how Medical Affairs strategies are implemented and executed within the umbrella of the overall product life cycle, interacting with various departments such as Marketing, Regulatory Affairs, Health Economics and Outcomes Research (HEOR)/Real World Evidence (RWE), and Clinical Development. The uniqueness of the UCB Medical Affairs – Immunology Fellowship is that it allows for optional rotations in the second year to various roles within Medical Affairs such as Medical Information, Medical Communications and Field Medical Operations & Strategy. This flexibility in the second year of the program allows for Fellows to gain broad experiences that will develop them into a well-rounded Medical Affairs professional.

Rotation	Timeframe
Medical Affairs Strategy – Immunology	1st Year (15 Months)
 Continue Medical Affairs Strategy Medical Information Medical Review Medical Digital Strategy Field Medical and Operations 	2nd Year (9 Months) Choices of 3-month interval rotations (Up to three)

Essential Functions & Responsibilities

- Gain scientific expertise in assigned disease areas within immunology to lead scientific and strategic discussions with key internal and external stakeholders
- Engage in key medical strategy tactics, including thought leader interactions, advisory board discussions, and align with the various immunology partners for portfolio and cross-therapeutic strategy
- Lead the execution of immunology medical deliverables including proactive patient management materials, medical proactive/ reactive decks, and training materials for cross-functional partners
- Participate in medical brand planning processes while representing the medical organization in cross-functional alignment calls
- Provide fair-balanced scientific responses to unsolicited requests from healthcare professionals regarding UCB products and help in the creation of Standardized Response Letters
- Assess ϑ identify gaps in MSL resources and collaborate with medical strategy on the development of MSL scientific resources and trainings
- Partner with the Field Medical Leadership Team to support development and implementation of field medical priorities
- Contribute to scientific congress Field Medical initiatives and engagement strategies
- Learn to conduct medical review of promotional and non-promotional materials in collaboration with Legal, Regulatory, and Marketing teams
- Gain an understanding of how HEOR contributes to the value of UCB products through real-world evidence and communication of value propositions to internal and external stakeholders



"A standout feature of the UCB Medical Affairs – Immunology Fellowship is the emphasis on Medical Affairs strategy. This allows Fellows to gain a well-round understanding of Medical Affairs, while allowing them to cultivate a robust skill set under the support and mentorship of experienced leadership. Additionally, the flexibility of the program empowers Fellows to directly engage in specific areas of interest allowing for further professional development. The foundation laid by this fellowship paves the way for Fellows to achieve a successful career as a Medical Affairs professional."

- Shelby Matthews, MA 1st Year Fellow

"UCB and the Medical Affairs Fellowship stood out to me for its focus on developing the Fellow in the strategy and execution of tactics of the medical team at a company that has a strong, patient-centered focus and a commitment to solving problems for patients with unmet needs. The rotational aspect of this fellowship allows the Fellow a breadth of experiences, while being a mid-size pharma company allows Fellows with the depth of experience I desired for my career development. Throughout the fellowship I feel that I have been supported and encouraged in every aspect to ensure that upon completion, I can be a complete, competitive, and successful Medical Affairs professional."



Global Clinical Sciences & Operations Fellowship

Recruiting 2 Fellows

Global Clinical Sciences & Operations (GCSO) at UCB aspires to deliver UCB's pipeline projects with top in the industry cycle times, strong patient focus, and innovative technologies. From efficient planning, to delivery of clinical trials of assigned products, to successful regulatory submissions for approval, GCSO is focused on the successful delivery of patient-preferred studies, thereby bringing value to our study participants and the global community.

The GCSO Fellowship will provide the Fellow with the unique opportunity to acquire in-depth end-to-end knowledge of the fundamentals of clinical trial operations. The 2-year Fellowship is designed to provide the Fellow with a variety of engaging rotational experiences to grow their knowledge and understanding of the many cross-functional teams required to execute highly rigorous clinical studies with high quality, within ambitious timelines, and ultimately providing value to our patients.

From planning, to execution, to reporting of clinical trials, the Fellow will gain first-hand experience rotating through the multidisciplinary sub-functions of GCSO including Clinical Project Management, Clinical Data & Innovation, Medical Writing, and Strategic Clinical Partnering.

Rotation	Timeframe
Clinical Project Management	5 months
Clinical Data & Innovation	5 months
Medical Writing	5 months
Strategic Clinical Partnering	5 months
Elective Rotation (outside GCSO)	2 months
Final Core Rotation (within GCSO)	3 months

Essential Functions & Responsibilities

- Develop an understanding of the end-to-end processes that enable execution of clinical trials
- Learn the various roles and responsibilities that contribute to clinical trial operations
- Participate in protocol design and protocol development
- Participate in logistical activities of study start-up such as supporting initial site feasibility, investigator selection, patient recruitment and engagement, site and vendor contracting, and study budget development
- Provide daily management of clinical trials, including interacting with external vendors including contract research organizations, technology vendors, and other third-party suppliers
- Leverage various digital platforms to perform clinical and data management activities
- Contribute to the adoption and incorporation of data innovation and various digital tools to drive diversity, equity, and inclusion in our trials
- Participate in study close-out activities including preparation of the clinical study report
- Gain experience in developing ICH compliant clinical and regulatory submission documents in support of drug approvals
- Have the opportunity to rotate through and interact with various sub-functions within GCSO such as: Patient Engagement, Feasibility, Risk Management, Data Standards, Trial Diversity
- Develop relationships and an extensive network within a cross-cultural, global organization, both within GCSO and with our stakeholders throughout the organization



"The GSCO fellowship program plays a pivotal role in ensuring the successful delivery of patient-centric clinical trials. Through hands-on experience across the entire clinical trial lifecycle, GSCO Fellows develop robust competencies that prepare them to become adaptable and well-rounded leaders in their field. Moreover, the program offers abundant opportunities for leadership and professional development within a supportive environment."

- Taysir A. Chamem, GCSO 1st Year Fellow

"UCB exemplifies the perfect blend of visionary innovation and unwavering commitment to patient-centricity. The Global Clinical Sciences and Operations (GCSO) fellowship at UCB is unique as it provides a nurturing environment that fosters hands-on learning, collaboration and empowers Fellows to become well rounded scientists. In addition to acquiring invaluable skills, Fellows are provided the opportunities to be forefront of the end-to-end processes involved in designing, executing, closing out and reporting of groundbreaking clinical studies."



Global Medical Affairs Fellowship

Not Recruiting for 2025–2027 Fellowship Term

The Global Medical Affairs Fellowship focuses on opportunities to learn, experience, and lead various activities involved in the dynamic functions of a global medical affairs organization. Fellows will utilize their time to learn how medical affairs strategies are developed, implemented, and executed across the different indications that make up the global medical team. Fellows will learn to work collaboratively and cross-functionally across regions while considering global needs. In the first 18 months, Fellows should expect to spend most of their time working in global medical, regional medical, and global medical communications including global content development, congress planning and execution, omnichannel and social media activities. Fellows will also have the opportunity to select an elective rotation opportunity of their choice within the Medical Affairs.

The uniqueness of the UCB Global Medical Affairs Fellowship is that it allows the Fellow the opportunity to learn and work in a truly global environment. Working in global medical affairs is uniquely rewarding, involving international collaboration to address medical challenges, bridge research, and real-world practices, and ensure global access to cutting-edge treatments. The Fellow can expect to engage with diverse stakeholders, gaining insights into various healthcare systems and regulatory landscapes, requiring adaptability and deep medical expertise. It's a one-of-a-kind opportunity to make a positive impact on global healthcare. Additionally, the option to select an elective within or outside of medical affairs will allow the Fellow to learn about another area of interest.

Rotation	Timeframe
Introduction to Medical Affairs (2 weeks) Therapeutic Area Training/Onboarding (2 weeks) Global Medical	12 months
US Medical Affairs	6 months
Elective Rotation	3 months
Global Medical Affairs (final rotation)	3 months

Essential Functions & Responsibilities

- Gain scientific expertise in assigned disease areas within global medical affairs to participate in scientific discussions with key internal and external stakeholders
- Engage in key global medical tactics, including global thought leader engagement, advisory board discussions, medical communications, and align with the various cross-functional partners
- Lead the execution of medical deliverables including proactive patient education materials, medical proactive/reactive decks, and support materials for cross-functional partners
- Provide support to the global medical information team to develop fair-balanced scientific global response documents (GRDs) to ultimately support the regions in the creation of Standardized Response Letters (SRLs)
- Assess & identify gaps in MSL resources and collaborate with medical communications strategy on the development of MSL scientific resources and training
- Partner with the Field Medical Leadership Team to support the development and implementation of field medical priorities
- Contribute to and lead parts of the global scientific congress planning and execution
- Learn to conduct medical reviews of promotional and non-promotional materials in collaboration with Legal, Regulatory, and Marketing teams
- Support the global medical communications with omnichannel activities including the creation of cutting-edge HCP educational content for social media etc.



"The Global Medical Affairs Fellowship Program helps you become an expert, develop invaluable skills, and gain experience in medical affairs on both a global and local scale. It provides great visibility to leadership, access to opportunities to gain insight on additional functional areas, and strong support for personal and professional growth. My experience with the program has been incredibly rewarding, and I truly believe it's shaping me into a leader in the field."

- Joelle Odigie, GMA 1st Year Fellow

"The Global Medical Affairs fellowship is an extraordinary opportunity, offering a truly global learning, and experiential, opportunity. Fellows will engage in direct, hands-on learning alongside esteemed leaders within the global and US medical affairs organization. It's a rare chance for pharmaceutical industry professionals to work in a global environment, while also benefiting from exposure to multiple unique assets. This fellowship presents a remarkable platform to gain invaluable experience, foster global collaboration, and ultimately make a meaningful impact on the lives of patients living with severe diseases. Upon completion, Fellows will possess a strong foundation, propelling them toward a successful career in medical affairs."



- Chioma Ezenduka, Global Medical Neuroimmunology Lead, UCB GMA Fellowship Program Director



"The UCB Global Clinical Sciences & Operations Fellowship will allow Fellows to dive into the world of clinical trial operations gaining first-hand experience and leveraging the insight and expertise of operations leaders throughout the organization. Not often in one's career are you provided the opportunity to touch activities from the design of a clinical trial all the way through to drug approval, so this Fellowship provides a unique end-to-end experience. We aim to provide a robust opportunity that once completed, will enable the Fellow to pursue a career in any number of fields related to clinical trial operations and execution."

- Amber Barnes, Head of Global Medical Writing, UCB GCSO Fellowship Program Director

"The Medical Safety and Pharmacovigilance Fellowship is a comprehensive program designed to equip participants with a range of skills in the field of product safety and pharmacovigilance for both drugs and medical devices. It offers an immersive experience that spans the entire product lifecycle (from development through post-marketing commitments) and provides a unique blend of scientific knowledge and practical operations. The fellowship not only focuses on enhancing technical expertise but also emphasizes leadership development, aiming to develop the next generation of individuals who will lead and innovate in the field of patient safety."



- Bella Sessoms, Head of Strategic Planning & Partnerships, UCB MS&PV Fellowship Program Director



"The UCB Medical Affairs Fellowship will allow Fellows to build a competitive skillset needed to succeed in various Medical Affairs functions, starting with strategy and then the execution of medical tactics. This unique program has the flexibility to allow Fellows to deep dive into specific functions that they are interested in with the support and mentorship of experienced leaders with the ultimate goal of preparing the Fellow to lead a successful career in Medical Affairs."

- **Tae Oh,** Senior Medical Solutions Lead Dermatology, UCB MA Fellowship Program Co-Director

"It is an exciting time to be a part of the UCB Medical Affairs Immunology team. As a UCB Medical Affairs Fellow, you will have many opportunities to collaborate across functions that meaningfully impact the scientific community. Here at UCB, we are committed to fostering a dynamic and supportive environment where innovation and patients are at the forefront."



- Cori Cooper, Senior Medical Solutions Lead Rheumatology, UCB MA Fellowship Program Co-Director

Application Process

Fellows will be selected on a nationally competitive basis, and candidates must have a Doctor of Pharmacy degree from an ACPEaccredited college of pharmacy by June 30, 2025. The Fellowship offers a competitive salary and benefits package.

Requirements

- Doctor of Pharmacy degree (PharmD)
- Graduate of an accredited and nationally recognized pharmacy school
- U.S. citizen or permanent resident

Qualifications

- Ability to work independently and proactively
- Ability to work in a collaborative, cross-cultural team environment and build effective partnerships
- Flexible and adaptable, and ability to work under pressure
- Excellent written and verbal communication skills knows when and how to communicate, using strong interpersonal skills and written communications when appropriate
- Analytical logically breaking situations or issues down into their essential elements: carrying out diagnosis and developing solutions
- Strong organizational and project management skills with a high level of attention to detail and time management skills
- Overriding commitment to integrity and high standards in self and others
- Able to understand and analyze clinical and medical data

How to Apply

This Fellowship position may only be applied for through the IPhO FellowMatch service: **FellowMatch | Industry Pharmacists Organization**

- A letter of intent and CV should be submitted through the FellowMatch portal.
- Two letters of recommendation should be submitted to **ucbpharmdfellowshipprogram@ucb.com**. In the subject line of the email, please enter the functional area acronym, followed by candidate's name.
- The application deadline is October 25th, 2024.
- Applications will be reviewed on a rolling basis, and applicants are encouraged to submit their materials on FellowMatch accordingly.

For questions regarding the Fellowship program, contact the Fellowship Director (below) or visit **U.S. PharmD Fellowships | UCB (ucb-usa.com)**

- Global Regulatory Affairs Fellowship (GRA): Iram Hasan (iram.hasan@ucb.com)
- Medical Safety and Pharmacovigilance Safety Fellowship (MS&PV): Bella Sessoms (bella.sessoms@ucb.com)
- Medical Affairs Immunology Fellowship (MA): Tae Oh (tae.oh@ucb.com) & Cori Cooper (cori.cooper@ucb.com)
- Global Clinical Sciences and Operations Fellowship (GCSO): Amber Barnes (amber.barnes@ucb.com)
- Global Medical Affairs Fellowship (GMA) Not Recruiting: Chioma Ezenduka (chioma.ezenduka@ucb.com)

Not Pictured: Taysir A. Chamem, GCSO 1st Year Fellow







UCB Atlanta Campus

The UCB Atlanta campus stands as a symbol of our longterm commitment to the Atlanta business community. Since opening our doors in 1994, this beautiful campus has grown from a handful of people to approximately 1,300 employees today. UCB is the largest biopharmaceutical company with a U.S. headquarters in the Atlanta area. We are conveniently located just a short drive from the heart of downtown Atlanta. Considered the capital of southern business, metro Atlanta is a thriving corporate hub which continues to attract top companies to the area, boosting the local economy and growing the population, which now exceeds 6 million people. Our proximity and easy access to Hartsfield-Jackson International Airport, one of the largest airports in the world, is key for UCB's global reach.

UCB RTP Campus

UCB has benefitted from a presence in the Research Triangle Park in North Carolina since 2001. The site in Morrisville is an integral part of UCB's vision to provide superior and sustainable value to patients with severe diseases. We are a dynamic workforce that is diversified with talented individuals, who bring a vibrant work environment and vitality to the RTP biotechnical area. From drug development, patient safety, and quality perspective, UCB employees continue to bring differentiated medicines to patients and physicians. UCB is proud to partner with academic institutions, like-minded businesses, as well as local and state government agencies. UCB has approximately 260 employees at its location in RTP. RTP is the largest and most prominent high-tech research and development park in the United States. Our proximity and easy access to Raleigh-Durham International Airport is key for UCB's global reach.



