Schrödinger offers powerful, easy to use solutions for CPG product research and development. Designed for non-expert and expert users alike, the Schrödinger platform offers simple workflows for building, simulating and analyzing real-life systems with advanced physics-based modeling and machine learning technology.

Schrödinger’s team of dedicated Scientific and Technology Support experts bring domain-level expertise to ensure users gain the most project impact from the platform. Additional research services and collaboration opportunities are available.

Here we present example applications of Schrödinger’s Consumer Packaged Goods capabilities.
Food and Beverage

Advance food processing and bio-based packaging
Predict water uptake, transport and plasticization in starch

Extend product shelf life
Predict chemical stability and understand antifungal mechanism of edible coating

Incorporate plant based-protein ingredients
Understand aggregation and emulsifying properties and mechanism for emulsions and foam

Predict and design flavors and scents
Understand taste mechanisms with deep learning and informatics-based methods

Understand food emulsions and architecture
Predict morphology of complex emulsions
Cosmetic and Personal Care

Optimize shampoo formulations
Model and quantify interface interactions between F-layers of hair follicles and shampoo formulations

Replace traditional surfactants
Simulate micelle formation, morphology, and self assembly of emulsifier into micelles

Understand microemulsions
Explore phase diagrams, multi-component separation/aggregation of microemulsion systems

Drive formulation development
Perform virtual screening of surfactant systems

Build realistic models for complex formulations
Construct mixtures of multiple ingredients, encapsulating polymers, and additives
Cleaning Products

**Improve cleaning efficiency**
Understand self assembly and morphology of cleaning surfactants on surfaces and the effect of environment (e.g. temperature) on the changes of efficiency

**Understand micelle formation**
Predict self assembly of surfactants

**Understand microemulsions**
Provide insight to structure, interfacial properties, and thermodynamics

**Advance formulation stability**
Predict stability, decomposition, and spectroscopic properties of molecules (e.g antioxidants)
Packaging

Develop sustainable packaging
Study interfacial interactions between packaging materials and consumer goods, and simulate water uptake for barrier design and performance

Minimize production waste
Predict thermomechanical properties of packaging materials

Innovate with natural materials
Explore active, recyclable, and bio-based materials through molecular simulation

Reduce processing cost
Investigate the ability to scale to manufacturing-level processes
Team Collaboration and Digital Data Management

Empower collaboration
Employ enterprise informatics tools for sharing experimental and predictive models seamlessly

Amplify research
Rapid deployment of machine learning models to drive predictions

Promote project management
Work side by side, accelerating project communication and collective learning; screen, share results, and make informed decisions
Products

Check on the products that enable your success in the CPG industry. Download our Materials Science Product Guide.

Desmond
Highly efficient Molecular Dynamics (MD) tool
- MD multi-stage workflow
- Free energy calculation (FEP+ solubility, metadynamics, replica exchange)
- Simulated annealing

AutoQSAR
Automated machine learning tool
- 1D, 2D and 3D descriptor builder
- Model building and auto-selection
- Make prediction

MS CG
MD tool with coarse-grained (CG)/mesoscopic sites
- Coarse-grained sketcher
- Coarse-grained force fields: Dissipative particle dynamics, Martini, Generalized Lennard-Jones + Coulomb

MS LiveDesign
Web-based enterprise informatics for materials design
- Digital data management
- Team collaborations across geographies and organizations

OPLS4
Advanced modern force field
- Force field builder
- Conformational analysis

MS Jaguar
Rapid ab initio electronic structure package
- Bond and ligand dissociation
- Reaction analysis
- Charge transport
Selected Publications

Molecular-Level Examination of Amorphous Solid Dispersion Dissolution
Mohammad Atif Faiz Afzal, Kristin Lehmkmper, Ekaterina Sobich, Thomas F. Hughes, David J. Giesen, Teng Zhang, Caroline M. Krauter, Paul Winget, Matthias Degenhardt, Samuel O. Kyeremateng*, Andrea R. Browning, and John C. Shelley*
* Mol. Pharmaceutics 2021, 18, 11, 3999-4014

The structural basis of odorant recognition in insect olfactory receptors
Josefin del Mármol, Mackenzie A. Yedlin & Vanessa Ruta
* Nature 2021, 597, 126-131

Characterizing moisture uptake and plasticization effects of water on amorphous amylose starch models using molecular dynamics methods
Jeffrey M.Sanders, Mayank Misra, Thomas J.L.Mustard, David J.Giesen, Teng Zhang, John Shelley, Mathew D.Halls
* Carbohydrate Polymers 2021, 252, 11716

Comprehensive structure-activity-relationship studies of sensory active compounds in licorice (Glycyrrhiza glabra)
Christian Schmid, Anne Brockhoff, Yaron Ben Shoshan-Galeczki, Maximilian Kranz, Timo D. Stark, Rukiye Erkaya, Wolfgang Meyerhof, Masha Y. Niv, Corinna Dawid, Thomas Hofmann
* Food Chemistry 2021, 364, 130420

In Silico Investigation of Bitter Hop-Derived Compounds and Their Cognate Bitter Taste Receptors
Andreas Dunkel, Thomas Hofmann, and Antonella Di Pizio
* J. Agric. Food Chem. 2020, 68, 38, 10414-10423

Structure-based screening for discovery of sweet compounds
Yaron Ben Shoshan-Galeczki, Masha Y Niv
* Food Chemistry 2020, 315, 126286

Bitter or not? BitterPredict, a tool for predicting taste from chemical structure
Dagan-Wiener, A.; Nissim, I.; Ben Abu, N.; Borgonovo, G.; Bassoli, A.; Niv, M.Y.
* Scientific Reports 2017, 7, 12074

Molecular Dynamics Simulation Study of Sodium Dodecyl Sulfate Micelle: Water Penetration and Sodium Dodecyl Sulfate Dissociation
Chun, B.J.; Choi, J.I.; Jang, S.S.
* Colloids and Surfaces A: Physicochemical and Engineering Aspects 2015, 474, 36