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Materials Genomics

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3 **Abstract**

4 This course introduces students to materials genomics, treating the periodic table and
5 all known crystal structures as a searchable, computable design space. Students learn
6 how materials databases are built, how to represent matter as numbers, graphs, or
7 fingerprints, how to interrogate and predict properties of solids, how to use ML as a
8 surrogate for quantum mechanics, and how to design new materials algorithmically.

9 **1 Course Information**

10 **4th Semester – 5 ECTS · 2h lecture + 2h exercises per week, together
11 with ML for Materials Processing & Characterization**

12 **2 Course Philosophy**

13 Materials genomics treats the periodic table and all known crystal structures as a
14 giant searchable, computable design space.

15 Students learn:

- 16 • how materials databases are built,
- 17 • how to represent matter as numbers, graphs, or fingerprints,
- 18 • how to interrogate and predict properties of solids,
- 19 • how to use ML as a surrogate for quantum mechanics,
- 20 • how to design new materials algorithmically.

21 **3 Week-by-Week Curriculum (14 weeks)**

22 **3.1 Unit I — Foundations of Materials Genomics (Weeks 1–3)**

23 **3.1.1 Week 1 – What is Materials Genomics?**

- 24 • Genomics analogy: genes → functions vs atoms → properties.
- 25 • Brief history: AFLOW, OQMD, Materials Project, NOMAD.
- 26 • PSPP from the structure-first viewpoint.

27 **Exercise:** Explore Materials Project; query bandgaps, energies, symmetries.

28 **3.1.2 Week 2 – Crystal structure fundamentals**

- 29 • Space groups, Wyckoff positions, symmetry operations.
- 30 • How symmetry informs descriptors.

31 **Exercise:** Using pymatgen / spglib to analyze symmetries.

32 **3.1.3 Week 3 – Materials databases & file formats**

- 33 • CIF, POSCAR, PDB-like formats.
- 34 • Thermodynamic quantities in databases: formation energy, stability, convex
35 hull.

36 **Exercise:** Parse CIF files, extract primitive cells, compute density.

37 **3.2 Unit II — Representations of Materials (Weeks 4–6)**

38 **3.2.1 Week 4 – Classical descriptors & materials fingerprints**

- 39 • Magpie, matminer.
- 40 • Stoichiometric, elemental, and structural features.

41 **Exercise:** Build a small property regressor with Magpie features.

42 **3.2.2 Week 5 – Graph-based representations**

- 43 • Crystal structures as graphs: nodes, edges, periodic boundary conditions.
- 44 • CGCNN, MEGNet architecture intuition (no training from scratch yet).

45 **Exercise:** Build a simple CGCNN-like graph featurizer.

46 **3.2.3 Week 6 – Local atomic environments**

- 47 • Voronoi tessellations, coordination numbers, SOAP descriptors.

- 48 • Role in interatomic potentials and ML force fields.

49 **Exercise:** Compute SOAP vectors; perform clustering in descriptor space.

50 **3.3 Unit III — High-Throughput Computation & Screening (Weeks 7–9)**

51 **3.3.1 Week 7 – Quantum mechanical data and DFT basics**

- 52 • What DFT gives you: energies, forces, band structures, elastic constants.
 53 • Why it's expensive; why ML surrogates matter.

54 **Exercise:** Run a toy DFT calculation (Quantum Espresso or MP workflows).

55 **3.3.2 Week 8 – High-throughput workflows**

- 56 • Automation: pymatgen, custodian, FireWorks, Atomate.
 57 • Data generation for building surrogate models.

58 **Exercise:** Perform a small FireWorks workflow (or simulate the idea without cluster
 59 resources).

60 **3.3.3 Week 9 – Phase stability & the convex hull**

- 61 • Formation energies, metastability, hull distance.
 62 • Mapping an entire chemical system.

63 **Exercise:** Reconstruct phase diagrams from Materials Project data.

64 **3.4 Unit IV — Learning Properties from Atomic Structure (Weeks 10–12)**

65 **3.4.1 Week 10 – Regression on crystal data**

- 66 • Predicting bandgaps, hardness, elastic moduli.
 67 • Comparing different representation families.

68 **Exercise:** Benchmark random forest, GPR, CGCNN on a small dataset.

69 **3.4.2 Week 11 – Machine-learned interatomic potentials**

- 70 • Overview: GAP, SNAP, MTP, NequIP.
 71 • Role in simulating defects, diffusion, mechanical behavior.

72 **Exercise:** Fit a tiny ML potential (ACE or simple SNAP-style) to toy data.

73 **3.4.3 Week 12 – Generative models for materials**

- 74 • VAEs, diffusion models for crystal generation.
 75 • Constraints: symmetry, stability, charge neutrality.

76 **Exercise:** Sample a generative model from a pretrained online source; analyze
 77 validity.

78 **3.5 Unit V — Mini-Project & Synthesis (Weeks 13–14)**

79 **3.5.1 Week 13 – Project workshop**

80 **Example projects:**

- 81 • Predict bandgap from composition + structure representation.
 82 • Identify new stable compounds in a chemical system.
 83 • Build a graph-based model for elastic constants.
 84 • Use ML to approximate formation energies for a ternary subsystem.
 85 • Analyze SOAP fingerprints across polymorphs.

86 **3.5.2 Week 14 – Presentations & Reflection**

- 87 • Interpreting models: SHAP for materials descriptors.
 88 • Strengths/limitations of materials genomics vs experiment-driven ML.
 89 • How computational and experimental ML meet in modern labs.

90 **4 Learning Outcomes**

91 Students completing this course will be able to:

- 92 • Navigate major materials databases and extract relevant structural/property
93 data.
- 94 • Represent crystals numerically using descriptors, fingerprints, and graphs.
- 95 • Train ML models to predict quantum-mechanical and thermodynamic proper-
96 ties.
- 97 • Analyze structural features via symmetry, coordination, and environments.
- 98 • Perform high-throughput screening of materials candidates.
- 99 • Understand and apply generative models for inorganic crystals.
- 100 • Critically evaluate ML results in computational materials discovery.