Psychometric analysis of LLM responses

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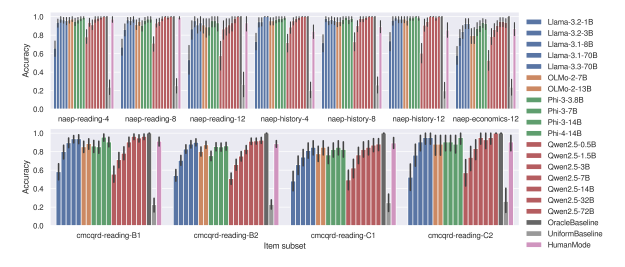
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Optimized KL divergence		
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import re		
from glob import glob		
from pathlib import Path		
<pre>import matplotlib.pyplot as plt</pre>		
import numpy as np		
import pandas as pd		
import scipy		
import seaborn as sns		
import torch		
from tqdm import tqdm		
<pre>sns.set_theme()</pre>		
<pre>plt.rcParams["font.family"] = "FreeSans"</pre>		
<pre>ITEMS_DIR = Path("/data/output")</pre>		
<pre>RESPONSES_DIR = Path("/llm-responses/output")</pre>		
<pre>FIGURES_DIR = Path("/paper/figures")</pre>		
MODELS = {		
"Llama-3.2-1B": "Llama",		
"Llama-3.2-3B": "Llama",		

```
"Llama-3.1-8B": "Llama",
    "Llama-3.1-70B": "Llama",
    "Llama-3.3-70B": "Llama",
    "OLMo-2-7B": "OLMo",
    "OLMo-2-13B": "OLMo",
    "Phi-3-3.8B": "Phi",
    "Phi-3-7B": "Phi",
    "Phi-3-14B": "Phi",
    "Phi-4-14B": "Phi",
    "Qwen2.5-0.5B": "Qwen",
    "Qwen2.5-1.5B": "Qwen",
    "Qwen2.5-3B": "Qwen",
    "Qwen2.5-7B": "Qwen",
    "Qwen2.5-14B": "Qwen",
    "Qwen2.5-32B": "Qwen",
    "Qwen2.5-72B": "Qwen",
    "OracleBaseline": "OracleBaseline",
    "UniformBaseline": "UniformBaseline",
    "HumanMode": "Human",
}
MODEL_FAMILY_COLORS = {
   "Llama": "#4c72b0",
    "OLMo": "#dd8452",
    "Phi": "#55a868",
    "Qwen": "#c44e52",
    "OracleBaseline": "#666666",
    "UniformBaseline": "#999999",
    "Human": "#da8bc3",
MODEL_PALETTE = [MODEL_FAMILY_COLORS[family] for model, family in MODELS.items()]
def load data(dataset, subject):
    items_df = pd.read_json(ITEMS_DIR / f"{dataset}-{subject}-items.jsonl", lines=True)
    response_distributions_df = pd.read_json(
        ITEMS_DIR / f"{dataset}-{subject}-response-distributions.jsonl", lines=True
    )
    responses_dfs = []
    for filename in glob(
        f"{dataset}-{subject}-responses-*.jsonl", root_dir=RESPONSES_DIR
    ):
```

```
(model,) = re.match(r".+-.+-responses-(.+)\.jsonl", filename).groups()
        df = pd.read_json(RESPONSES_DIR / filename, lines=True)
        df["model"] = model
        responses_dfs.append(df)
    responses_df = pd.concat(responses_dfs)
    responses_df["dataset"] = dataset
    responses_df["subject"] = subject
    responses_df = items_df.merge(response_distributions_df, on="item_id").merge(
        responses_df, on="item_id"
    )
    if dataset == "naep":
        responses_df.sort_values(["grade"], inplace=True)
        responses_df["subset"] = (
            responses_df[["dataset", "subject", "grade"]]
            .astype(str)
            .agg("-".join, axis=1)
    elif dataset == "cmcqrd":
        responses_df.sort_values(["level"], inplace=True)
        responses_df["subset"] = (
            responses_df[["dataset", "subject", "level"]]
            .astype(str)
            .agg("-".join, axis=1)
        )
    return responses_df
naep_data = pd.concat(
    Γ
        load_data(dataset, subject)
        for dataset, subject in [
            ("naep", "reading"),
            ("naep", "history"),
            ("naep", "economics"),
        ]
    ]
cmcqrd_data = load_data("cmcqrd", "reading")
```

```
for data in [naep_data, cmcqrd_data]:
    # Ensure consistent order of subsets and models
    subsets = data["subset"].unique()
    data["subset"] = data["subset"].astype("category").cat.set_categories(subsets)
    data["model"] = data["model"].astype("category").cat.set_categories([model for model in ])
    assert data["model"].notna().all(), "Missing model"
    data["model_family"] = data["model"].map(MODELS)
    data.sort_values(["subset", "model"], inplace=True)
    data.reset_index(drop=True, inplace=True)
    data["human_probs"] = data["response_distribution"].apply(
        lambda response_distribution: np.array(response_distribution)
        / sum(response_distribution)
    data["model_probs"] = data["logits"].apply(
        lambda logits: scipy.special.softmax(logits, axis=1).mean(axis=0)
    data["model_correct"] = data.apply(
        lambda row: np.argmax(row["model_probs"]) == row["correct_option_index"], axis=1
    )
# Accuracy
fig, (ax1, ax2) = plt.subplots(2, 1, figsize=(12, 5.5), sharey=True)
# Add HumanMode as a model
naep_accuracy_models = naep_data[["subset", "model", "model_correct"]].copy()
naep_accuracy_human = naep_data.groupby(["subset", "item_id"], observed=True).first().reset_
naep_accuracy_human["model"] = "HumanMode"
naep_accuracy_human["model_correct"] = naep_accuracy_human.apply(
    lambda row: np.argmax(row["response_distribution"]) == row["correct_option_index"], axis
naep_accuracy_all = pd.concat([naep_accuracy_models, naep_accuracy_human])
cmcqrd_accuracy_models = cmcqrd_data[["subset", "model", "model_correct"]].copy()
cmcqrd_accuracy_human = cmcqrd_data.groupby(["subset", "item_id"], observed=True).first().re
cmcqrd_accuracy_human["model"] = "HumanMode"
cmcqrd_accuracy_human["model_correct"] = cmcqrd_accuracy_human.apply(
    lambda row: np.argmax(row["response_distribution"]) == row["correct_option_index"], axis
cmcqrd_accuracy_all = pd.concat([cmcqrd_accuracy_models, cmcqrd_accuracy_human])
sns.barplot(
```

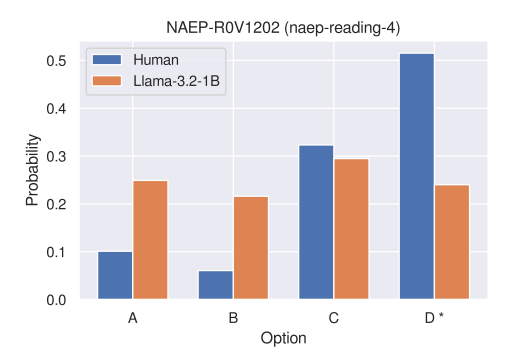
```
data=naep_accuracy_all,
    x="subset",
    y="model_correct",
    hue="model",
    width=0.9,
    palette=MODEL_PALETTE,
    err_kws={"linewidth": 1.5},
    ax=ax1,
ax1.set_xlabel("")
ax1.set_ylabel("Accuracy")
sns.barplot(
    data=cmcqrd_accuracy_all,
   x="subset",
    y="model_correct",
    hue="model",
    width=0.9,
    palette=MODEL_PALETTE,
    legend=False,
    ax=ax2,
ax2.set_xlabel("Item subset")
ax2.set_ylabel("Accuracy")
ax1.legend(loc="upper left", bbox_to_anchor=(1, 1), frameon=False)
fig.savefig(FIGURES_DIR / "accuracy.pdf", bbox_inches="tight")
```



```
def plot_probs(
    response,
    model_probs_col="model_probs",
   human_probs_col="human_probs",
    reference_prob_col=None,
    reference_prob_label="Reference",
):
   human_probs = response[human_probs_col]
   model_probs = response[model_probs_col]
   fig, ax = plt.subplots()
    bar_width = 0.35
    index = np.arange(len(human_probs))
    ax.bar(index, human_probs, bar_width, label="Human")
    ax.bar(index + bar_width, model_probs, bar_width, label=response["model"])
    if reference_prob_col:
        reference_prob = response[reference_prob_col]
        ax.plot(
            [-0.5, len(human_probs) - 0.5],
            [reference_prob, reference_prob],
            color="black",
            linestyle="--",
            label=reference_prob_label,
        )
    ax.set_xlabel("Option")
    ax.set_ylabel("Probability")
    ax.set_title(f"{response['item_id']} ({response['subset']})")
    ax.set_xticks(index + bar_width / 2)
    labels = ["A", "B", "C", "D"]
```

```
labels[response["correct_option_index"]] += " *"
ax.set_xticklabels(labels)
ax.legend()

plot_probs(naep_data.iloc[0])
```



Temperature optimization

We optimize the temperature separately for each item subset (subject and NAEP grade or CMCQRD level) to minimize the KL divergence between the LLM response distribution and the human response distribution.

NOTE: We optimize on the same data that we evaluate on, in order to get an optimistic estimate / upper bound of psychometric plausibility in the following analyses.

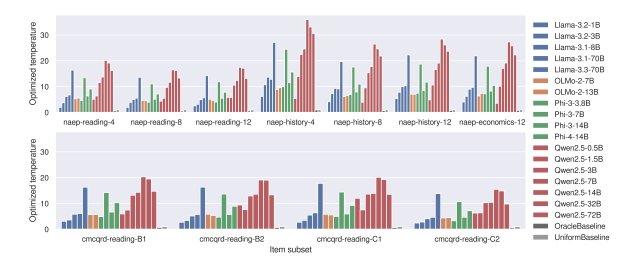
```
def optimize_temperature(logits, human_probs, init=1.0):
    logits = torch.tensor(list(logits))
    human_probs = torch.tensor(list(human_probs))
    temperature = torch.tensor(init, requires_grad=True)
```

```
optimizer = torch.optim.Adam([temperature], lr=0.01)
   best_loss = float("inf")
   steps_without_improvement = 0
   while True:
       optimizer.zero_grad()
       tempered_logits = logits / temperature
       model_logprobs = torch.nn.functional.log_softmax(tempered_logits, dim=-1)
       model_logprobs = model_logprobs.mean(dim=-2)
       loss = torch.nn.functional.kl_div(
           model_logprobs, human_probs.clone(), reduction="batchmean"
       loss.backward()
       optimizer.step()
       if loss.item() >= best_loss:
            steps_without_improvement += 1
            if steps_without_improvement > 100:
               break
        else:
            steps_without_improvement = 0
           best_loss = loss.item()
   return temperature.item()
def grouped_optimize_temperature(df):
   optimized_temperatures = pd.Series(index=df.index, dtype=float)
   for _, group in tqdm(
       df.groupby(["subset", "model"], observed=True), desc="Optimizing temperatures"
   ):
       optimized_temperatures[group.index] = optimize_temperature(
           group["logits"], group["human_probs"]
   return optimized_temperatures
for data in [naep_data, cmcqrd_data]:
   data["optimized_temperature"] = grouped_optimize_temperature(data)
```

```
data["optimized_model_probs"] = data.apply(
        lambda row: scipy.special.softmax(
            np.array([*row["logits"]]) / row["optimized_temperature"], axis=1
        ).mean(axis=0),
        axis=1,
                           0%1
                                         | 0/140 [00:00<?, ?it/s]/tmp/ipykernel_1473869/177978
Optimizing temperatures:
  human_probs = torch.tensor(list(human_probs))
                                         | 1/140 [00:02<05:49, 2.51s/it]Optimizing temperatu
Optimizing temperatures:
                           1%|
Optimizing temperatures:
                           0%|
                                         | 0/80 [00:00<?, ?it/s]Optimizing temperatures:
fig, (ax1, ax2) = plt.subplots(2, 1, figsize=(12, 5.5), sharey=True)
sns.barplot(
    data=naep_data,
    x="subset",
    y="optimized_temperature",
    hue="model",
    errorbar=None,
    width=0.9,
    palette=MODEL_PALETTE,
    ax=ax1,
ax1.set_xlabel("")
ax1.set_ylabel("Optimized temperature")
sns.barplot(
    data=cmcqrd_data,
    x="subset",
    y="optimized_temperature",
    hue="model",
    errorbar=None,
    width=0.9,
    palette=MODEL_PALETTE,
    legend=False,
    ax=ax2,
ax2.set_xlabel("Item subset")
ax2.set_ylabel("Optimized temperature")
```

```
ax1.legend(loc="upper left", bbox_to_anchor=(1, 1), frameon=False)
fig.savefig(FIGURES_DIR / "temperature.pdf", bbox_inches="tight")
```

```
/tmp/ipykernel_1473869/2049676019.py:3: UserWarning: The palette list has more values (21) to
sns.barplot(
/tmp/ipykernel_1473869/2049676019.py:16: UserWarning: The palette list has more values (21)
sns.barplot(
```



Response distribution analysis

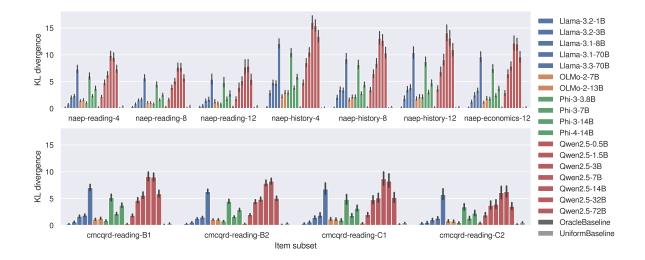
In this analysis, we compare the token probability distribution produced by the LLMs to the human response distribution.

```
for data in [naep_data, cmcqrd_data]:
    data["kldiv"] = data.apply(
        lambda row: scipy.stats.entropy(row["human_probs"], row["model_probs"]), axis=1
)
    data["optimized_kldiv"] = data.apply(
        lambda row: scipy.stats.entropy(
            row["human_probs"], row["optimized_model_probs"]
        ),
        axis=1,
)
```

Unoptimized KL divergence

```
fig, (ax1, ax2) = plt.subplots(2, 1, figsize=(12, 5.5), sharey=True)
sns.barplot(
    data=naep_data,
   x="subset",
    y="kldiv",
    hue="model",
    width=0.9,
    palette=MODEL_PALETTE,
    err_kws={"linewidth": 1.5},
    ax=ax1,
ax1.set_xlabel("")
ax1.set_ylabel("KL divergence")
sns.barplot(
    data=cmcqrd_data,
    x="subset",
    y="kldiv",
    hue="model",
    width=0.9,
    palette=MODEL_PALETTE,
    legend=False,
    ax=ax2,
)
ax2.set_xlabel("Item subset")
ax2.set_ylabel("KL divergence")
ax1.legend(loc="upper left", bbox_to_anchor=(1, 1), frameon=False)
fig.savefig(FIGURES_DIR / "kldiv-unoptimized.pdf", bbox_inches="tight")
```

```
/tmp/ipykernel_1473869/1437317739.py:3: UserWarning: The palette list has more values (21) to
    sns.barplot(
/tmp/ipykernel_1473869/1437317739.py:16: UserWarning: The palette list has more values (21) sns.barplot(
```



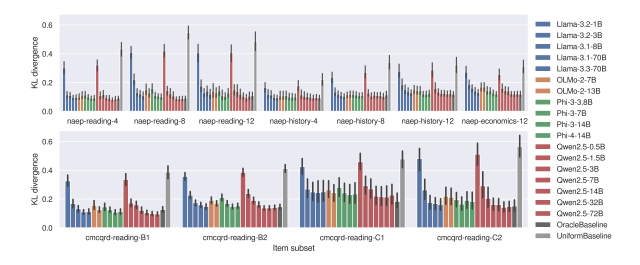
Optimized KL divergence

```
fig, (ax1, ax2) = plt.subplots(2, 1, figsize=(12, 5.5), sharey=True)
sns.barplot(
    data=naep_data,
    x="subset",
    y="optimized_kldiv",
    hue="model",
    width=0.9,
    palette=MODEL_PALETTE,
    err_kws={"linewidth": 1.5},
    ax=ax1,
ax1.set_xlabel("")
ax1.set_ylabel("KL divergence")
sns.barplot(
    data=cmcqrd_data,
    x="subset",
    y="optimized_kldiv",
    hue="model",
    width=0.9,
    palette=MODEL_PALETTE,
    legend=False,
    ax=ax2,
```

```
)
ax2.set_xlabel("Item subset")
ax2.set_ylabel("KL divergence")
ax1.legend(loc="upper left", bbox_to_anchor=(1, 1), frameon=False)
fig.savefig(FIGURES_DIR / "kldiv-optimized.pdf", bbox_inches="tight")
```

/tmp/ipykernel_1473869/3176962260.py:3: UserWarning: The palette list has more values (21) to
sns.barplot(
/tmp/ipykernel_1473869/3176962260.py:16: UserWarning: The palette list has more values (21)

/tmp/ipykernel_1473869/3176962260.py:16: UserWarning: The palette list has more values (21)
sns.barplot(



CTT analysis

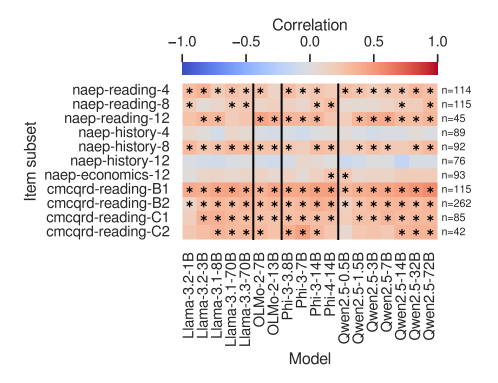
In this analysis, we compare the item facility values (i.e., the proportion of students who answered each item correctly) to the probability of the correct answer option produced by the LLMs. Psychometrically plausible LLM responses should be more confidently correct with easier items than with more difficult items.

```
# Baselines are not useful for comparison (same probability for all items)
naep_data = naep_data[~naep_data["model"].str.endswith("Baseline")]
cmcqrd_data = cmcqrd_data[~cmcqrd_data["model"].str.endswith("Baseline")]
for data in [naep_data, cmcqrd_data]:
```

```
data["model"] = data["model"].cat.remove_unused_categories()
    data["human_correct_prob"] = data.apply(
        lambda row: row["human_probs"][row["correct_option_index"]], axis=1
    )
    data["model_correct_prob"] = data.apply(
        lambda row: row["model_probs"][row["correct_option_index"]], axis=1
    )
    data["optimized_model_correct_prob"] = data.apply(
        lambda row: row["optimized model_probs"][row["correct_option_index"]], axis=1
    )
all_data = pd.concat([naep_data, cmcqrd_data])
all_data["subset"] = (
    all_data["subset"]
    .astype("category")
    .cat.set_categories(
        naep_data["subset"].cat.categories.append(cmcqrd_data["subset"].cat.categories)
# Correlation between LLM and human response probabilities
correct_probs_human_corr = (
    all_data.groupby(["subset", "model"], observed=True)
    .apply(
        lambda group: scipy.stats.pearsonr(
            group["human_correct_prob"],
            group["optimized_model_correct_prob"],
            alternative="two-sided",
    )
    .apply(pd.Series, index=["correlation", "p-value"])
).unstack()
# Heatmap of correlations
fig, ax = plt.subplots(figsize=(7.0, 4.0))
sns.heatmap(
    correct_probs_human_corr["correlation"],
    annot=(correct_probs_human_corr["p-value"] < 0.05).replace({True: "*", False: ""}),</pre>
    annot_kws={"color": "black"},
    fmt="",
    cmap="coolwarm",
```

```
center=0,
    vmin=-1,
    vmax=1,
    cbar_kws={"label": "Correlation", "location": "top", "fraction": 0.072},
    square=True,
    ax=ax,
ax.set_xlabel("Model")
ax.set_ylabel("Item subset")
# Lines between model families
for i, (model, next_model) in enumerate(
    zip(
        correct_probs_human_corr["correlation"].columns[:-1],
        correct_probs_human_corr["correlation"].columns[1:],
):
    if MODELS.get(model) != MODELS.get(next_model):
        ax.axvline(i + 1, color="black")
# Sample size per subset, and model
sample_size = (
    all_data.groupby(["subset", "model"], observed=True).size().unstack().iloc[:, 0]
for i, index in enumerate(correct_probs_human_corr.index):
    ax.annotate(
        f"n={sample_size[index]:.0f}",
        xy=(len(correct_probs_human_corr["correlation"].columns) + 0.3, i + 0.5),
        ha="left",
        va="center",
        annotation_clip=False,
        size=8,
    )
fig.tight_layout()
fig.savefig(FIGURES_DIR / "ctt-correlation.pdf", bbox_inches="tight")
```

/tmp/ipykernel_1473869/2245631322.py:4: DeprecationWarning: DataFrameGroupBy.apply operated
.apply(



Subset with strongest correlation
strongest_subset = correct_probs_human_corr["correlation"].max(axis="columns").idxmax()
correct_probs_human_corr.loc[strongest_subset, "correlation"]

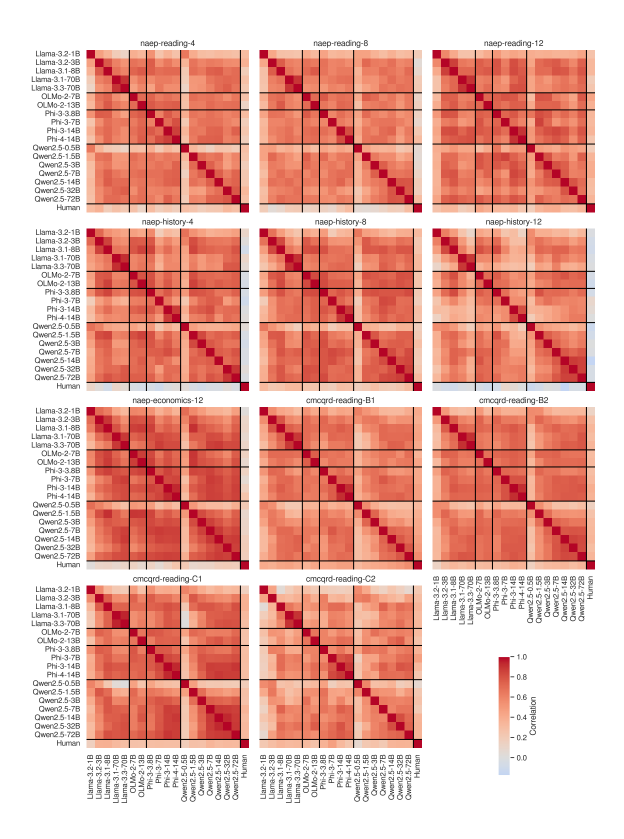
model	
Llama-3.2-1B	0.408693
Llama-3.2-3B	0.388440
Llama-3.1-8B	0.439477
Llama-3.1-70B	0.457404
Llama-3.3-70B	0.412323
OLMo-2-7B	0.366660
OLMo-2-13B	0.432958
Phi-3-3.8B	0.422561
Phi-3-7B	0.424779
Phi-3-14B	0.465331
Phi-4-14B	0.465496
Qwen2.5-0.5B	0.323238
Qwen2.5-1.5B	0.386761
Qwen2.5-3B	0.419301
Qwen2.5-7B	0.432884
Qwen2.5-14B	0.472846

```
Qwen2.5-72B
                 0.557089
Name: cmcqrd-reading-B1, dtype: float64
# Full correlation matrices for all LLMs and human
correct probs = all data.pivot(
    index=["item_id", "subset"], columns="model", values="optimized_model_correct_prob"
correct_probs["Human"] = all_data.groupby(["item_id", "subset"])[
    "human_correct_prob"
1.first()
correct_probs_corr = correct_probs.groupby("subset").corr()
g = sns.FacetGrid(
    correct_probs_corr.reset_index("subset"), col="subset", col_wrap=3, height=4
def heatmap(data, **kwargs):
    data = data.drop(columns="subset")
    sns.heatmap(data, **kwargs)
g.map_dataframe(heatmap, cmap="coolwarm", center=0, cbar=False, square=True)
g.set_titles("{col_name}")
# Color bar
cbar_ax = g.figure.add_axes([0.82, 0.06, 0.02, 0.15])
g.figure.colorbar(g.facet_axis(0, 5).collections[0], cax=cbar_ax, label="Correlation")
# Lines between model families
for ax in g.axes:
   ax.set_xlabel("")
   ax.set_ylabel("")
    for i, (model, next_model) in enumerate(
        zip(correct_probs_corr.columns[:-1], correct_probs_corr.columns[1:])
    ):
        if MODELS.get(model) != MODELS.get(next_model):
            ax.axvline(i + 1, color="black")
            ax.axhline(i + 1, color="black")
g.figure.subplots_adjust(hspace=0.1)
g.savefig(FIGURES_DIR / "ctt-correlation-full.pdf", bbox_inches="tight")
```

Qwen2.5-32B

0.508665

```
/tmp/ipykernel_1473869/1301630354.py:5: FutureWarning: The default of observed=False is depredered correct_probs["Human"] = all_data.groupby(["item_id", "subset"])[
/tmp/ipykernel_1473869/1301630354.py:8: FutureWarning: The default of observed=False is depredered correct_probs_corr = correct_probs.groupby("subset").corr()
```



IRT analysis

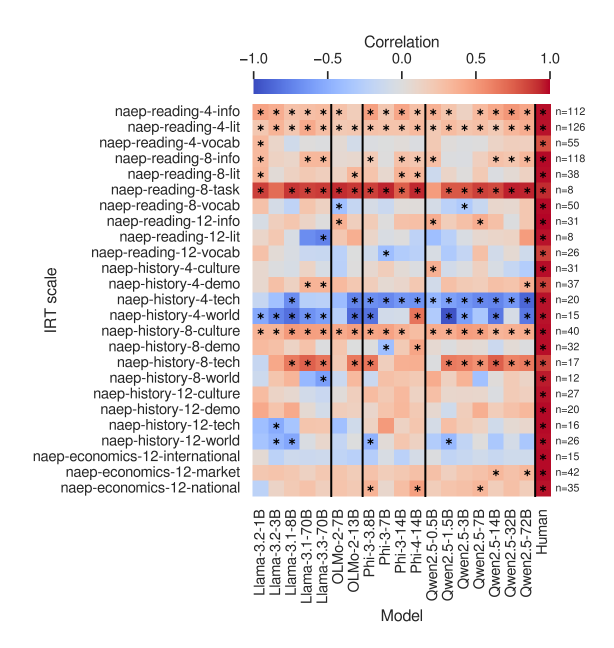
In this analysis, we study how well the LLM response probabilities fit a three-parameter logistic IRT model fitted to human responses. Specifically, we view each LLM as a single test taker with $\theta = 0$ (i.e., average ability) and check whether this test taker's response probabilities correlate with the expected probabilities from the IRT model.

NOTE: This analysis is only applicable to NAEP data, as CMCQRD data only contains rescaled difficulty parameters, meaning that we cannot reconstruct the response probability for the average test taker.

```
naep_parameters_df = pd.read_json(ITEMS_DIR / "naep-parameters.json1", lines=True)
naep_parameters_df = naep_parameters_df [naep_parameters_df ["irt_model"] == "3pl"].drop(
    columns=["irt_model", "d_1", "d_2", "d_3", "d_4"]
)
naep_irt_data = naep_data.merge(naep_parameters_df, on=["item_id", "subject", "grade"])
# Expected probability of correct response at theta = 0
def p_at_theta0(a, b, c):
    return c + (1 - c) / (1 + np.exp(-1.7 * a * (0 - b)))
naep_irt_data["p_at_theta0"] = naep_irt_data.apply(
    lambda row: p_at_theta0(row["a"], row["b"], row["c"]), axis=1
)
# Correlation between LLM response probabilities and IRT model probabilities
correct probs p corr = (
    naep_irt_data.groupby(["subset", "scale", "model"], observed=True)
    .apply(
        lambda group: scipy.stats.pearsonr(
            group["p_at_theta0"],
            group["optimized_model_correct_prob"],
            alternative="two-sided",
        ),
        include groups=False,
    .apply(pd.Series, index=["correlation", "p-value"])
).unstack()
# Correlation between human response probabilities and IRT model probabilities
correct_probs_p_human_corr = (
```

```
naep_irt_data
    # Deduplicate response distributions (keep only one model)
    .groupby(["subset", "scale", "item_id", "year"], observed=True)
    .first()
    # Correlate
    .groupby(["subset", "scale"], observed=True)
        lambda group: scipy.stats.pearsonr(
            group["p_at_theta0"],
            group["human_correct_prob"],
            alternative="two-sided",
        ),
        include_groups=False,
    .apply(pd.Series, index=["correlation", "p-value"])
correct_probs_p_corr["correlation", "Human"] = correct_probs_p_human_corr["correlation"]
correct_probs_p_corr["p-value", "Human"] = correct_probs_p_human_corr["p-value"]
# Heatmap of correlations
fig, ax = plt.subplots(figsize=(7.5, 6.5))
g = sns.heatmap(
    correct_probs_p_corr["correlation"],
    annot=(correct_probs_p_corr["p-value"] < 0.05).replace({True: "*", False: ""}),</pre>
    annot kws={"color": "black"},
   fmt="",
    cmap="coolwarm",
   center=0,
   vmin=-1,
   vmax=1,
    cbar_kws={"label": "Correlation", "location": "top", "fraction": 0.0355, "pad": 0.025},
    square=True,
    ax=ax,
)
ax.set_xlabel("Model")
ax.set_ylabel("IRT scale")
ax.yaxis.labelpad = -20
ax.set_yticklabels(
    f"{subset}-{scale}" if scale else subset
        for subset, scale in correct_probs_p_corr.index
    ]
```

```
# Lines between model families
for i, (model, next_model) in enumerate(
    zip(
        correct_probs_p_corr["correlation"].columns[:-1],
        correct_probs_p_corr["correlation"].columns[1:],
    )
):
    if MODELS.get(model) != MODELS.get(next_model):
        ax.axvline(i + 1, color="black")
# Sample size per subset, scale, and model
sample_size = (
    naep_irt_data.groupby(["subset", "scale", "model"], observed=True)
    .size()
    .unstack()
    .iloc[:, 0]
for i, index in enumerate(correct_probs_p_corr.index):
    ax.annotate(
        f"n={sample_size[index]:.0f}",
        xy=(len(correct_probs_p_corr["correlation"].columns) + 0.3, i + 0.5),
        ha="left",
        va="center",
        annotation_clip=False,
        size=8,
    )
fig.tight_layout()
fig.savefig(FIGURES_DIR / "irt-correlation.pdf", bbox_inches="tight")
```



```
# Regression plots of LLM response probabilities vs. IRT model probabilities
naep_irt_data["subject-scale"] = naep_irt_data["subject"] + "-" + naep_irt_data["scale"]
g = sns.lmplot(
    naep_irt_data,
    x="p_at_theta0",
    y="optimized_model_correct_prob",
    hue="model",
    col="subject-scale",
```

```
row="grade",
)
g.set(xlim=(0, 1), ylim=(0, 1))
g.set_titles("NAEP {col_name}, grade {row_name}")
g.set_axis_labels(
    "Expected probability at $\\theta = 0$", "Observed response probability"
)
g.savefig(FIGURES_DIR / "irt-regression.pdf", bbox_inches="tight")
```

