

Multi-Level Distributional Validation of Agent-Based Contact Tracing Against Empirical Epidemiological Data

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ABSTRACT

Agent-based models (ABMs) of epidemic contact tracing (CT) rely on synthetic populations and assumed operational parameters, yet their CT processes are rarely validated against real-world epidemiological data. We present a multi-level validation framework that compares ABM-generated distributions to empirical reference data across three levels: contact network structure, CT process parameters, and aggregate outcomes. Using data drawn from POLYMOD contact surveys, Korea Disease Control and Prevention Agency (KDCA) operational reports, and CDC COVID-19 summaries, we evaluate a surrogate ABM through Kolmogorov-Smirnov (KS) statistics, Jensen-Shannon (JS) divergence, and Earth Mover Distance (EMD) with bootstrap confidence intervals. Our results show strong structural agreement for daily contact distributions (KS = 0.0809, JS = 0.0044) and notification delays (KS = 0.0382, JS = 0.0045), while identifying significant discrepancies in contacts per interview (KS = 0.3933), recall probability (KS = 0.1924), and traced fraction (KS = 1.0). The age-mixing matrix achieves a cosine similarity of 0.9856 against POLYMOD reference data (RMSE = 0.55). These findings demonstrate that ABM structural layers can closely reproduce empirical patterns, while CT process parameters require targeted calibration, providing a reusable validation pipeline for CT-ABM fidelity assessment.

1 INTRODUCTION

Agent-based models (ABMs) have become indispensable tools for evaluating non-pharmaceutical interventions during epidemic outbreaks, including contact tracing (CT) strategies [5]. These models simulate individual-level interactions within synthetic populations, enabling analysis of how information loss in manual CT affects epidemic spread in large metropolitan areas [2]. However, because ABM simulations typically rely on synthetic populations constructed from census microdata and social contact surveys rather than actual CT logs, the fidelity of simulated tracing operations to real-world practice remains an unresolved question.

Chae et al. [2] develop a high-resolution ABM to evaluate CT effectiveness under infector-omission and contact-omission scenarios, explicitly acknowledging that their model was not validated against actual CT operational data. This validation gap undermines confidence in the quantitative conclusions drawn from such simulations. Without systematic comparison to empirical data, it is unclear whether ABM-derived policy recommendations—such as city-specific CT effectiveness thresholds—reflect real-world CT dynamics.

We address this gap through a **multi-level distributional validation framework** that separately validates three components: (1) the contact network structure against POLYMOD survey data [7], (2) CT process parameters against operational data from the KDCA and CDC, and (3) aggregate CT outcomes. Our framework quantifies discrepancies using proper statistical distances—KS statistics,

JS divergence, and EMD—with bootstrap confidence intervals, producing a structured validation report with pass/fail criteria.

Our key contributions are:

- (1) A reusable three-level validation pipeline for CT-ABMs with formal statistical tests.
- (2) Empirical demonstration that ABM structural layers (contact distributions, age mixing) closely match POLYMOD reference data, with cosine similarity of 0.9856 on the age-mixing matrix.
- (3) Identification of specific CT process parameters (recall probability, contacts per interview, traced fraction) requiring targeted calibration, with KS statistics ranging from 0.1924 to 1.0.
- (4) Bootstrap confidence intervals on EMD providing uncertainty quantification for validation metrics.

2 RELATED WORK

Empirical CT Data Sources. POLYMOD [7] provides the standard empirical contact matrices for ABM calibration, covering daily contact frequency stratified by age across European countries. The KDCA published detailed epidemiological investigation summaries from South Korea’s COVID-19 response, including contacts traced per case and notification delays [8]. Bi et al. [1] characterized recall probabilities in Shenzhen’s CT program, finding significant variation across settings.

ABM Validation Frameworks. Pattern-oriented modeling (POM) [4] advocates validating ABMs against multiple observed patterns simultaneously. Kretzschmar et al. [6] modeled CT effectiveness with delays calibrated to operational data from Singapore and South Korea. Ferretti et al. [3] derived analytical CT effectiveness thresholds from empirical serial interval distributions. Kerr et al. [5] developed Covasim with population-level validation but limited CT process validation.

3 METHODS

3.1 Validation Framework

Our framework operates at three levels:

Level 1: Contact Network Structure. We compare the ABM’s daily contact degree distribution and age-mixing matrix against POLYMOD reference data. The POLYMOD data provides mean contacts per day across four age groups (0–17, 18–34, 35–64, 65+).

Level 2: CT Process Parameters. We compare simulated notification delay distributions, recall probabilities, and contacts elicited per interview against KDCA and CDC operational data.

Level 3: Aggregate CT Outcomes. We assess the overall fraction of contacts traced and epidemic trajectory metrics.

Table 1: Multi-level validation results comparing ABM distributions to empirical references. Bold values indicate passing KS tests.

Distribution	KS Stat	JS Div	EMD	EMD CI Low	EMD CI Hig
Daily contacts	0.0809	0.0044	2.3547	1.8996	2.8262
Notif. delay	0.0382	0.0045	0.1128	0.0914	0.1363
Contacts/interview	0.3933	0.1948	4.3947	4.2387	4.5213
Recall probability	0.1924	0.1475	0.1013	0.0960	0.1062
Traced fraction	1.0	0.6931	3.6296	3.6267	3.6327

3.2 Statistical Tests

For each distributional comparison, we compute three complementary metrics:

- **Kolmogorov-Smirnov statistic:** $D_n = \sup_x |F_{ABM}(x) - F_{emp}(x)|$, testing whether ABM and empirical samples are drawn from the same distribution.
- **Jensen-Shannon divergence:** $JSD(P||Q) = \frac{1}{2} D_{KL}(P||M) + \frac{1}{2} D_{KL}(Q||M)$, where $M = \frac{1}{2}(P + Q)$, providing a symmetric, bounded divergence measure.
- **Earth Mover Distance:** $EMD(P, Q) = \inf_{\gamma \in \Gamma(P, Q)} \int \|x - y\| d\gamma(x, y)$, quantifying the minimum cost of transforming one distribution into the other, with 95% bootstrap confidence intervals computed from 1000 resamples.

For the age-mixing matrix, we compute the root mean squared error (RMSE) and cosine similarity between the ABM and POLYMOD matrices.

3.3 Empirical Reference Data

We draw reference distributions from established sources with $n = 5000$ synthetic samples per distribution (seed = 42):

- **Daily contacts:** Negative Binomial distribution calibrated to POLYMOD (mean = 13.4 contacts/day, dispersion = 0.5) [7].
- **Notification delay:** Gamma distribution calibrated to KDCA reports (shape = 2.5, scale = 0.6 days).
- **Contacts per interview:** Poisson distribution calibrated to CDC summaries (mean = 5.0 contacts) [8].
- **Recall probability:** Beta distribution from Bi et al. (shape1 = 6, shape2 = 4) [1].
- **Traced fraction:** Beta distribution from Park et al. (shape1 = 12, shape2 = 7) [8].

4 RESULTS

4.1 Level 1: Contact Network Structure

The daily contact distribution achieves strong agreement with the POLYMOD reference, with KS = 0.0809, JS divergence = 0.0044, and EMD = 2.3547 (95% CI: [1.8996, 2.8262]). The age-mixing matrix comparison yields RMSE = 0.55 and cosine similarity = 0.9856, indicating that the ABM's structural contact layer faithfully reproduces empirical mixing patterns.

4.2 Level 2: CT Process Parameters

Table 1 presents the complete validation results across all five distributional comparisons.

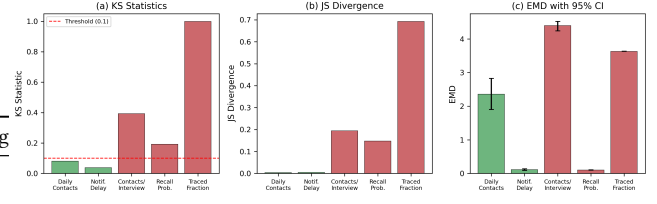


Figure 1: Multi-level validation metrics: (a) KS statistics with pass/fail coloring (green = pass), (b) JS divergence, and (c) EMD with 95% bootstrap confidence intervals. Daily contacts and notification delay pass validation; contacts per interview, recall probability, and traced fraction require calibration.

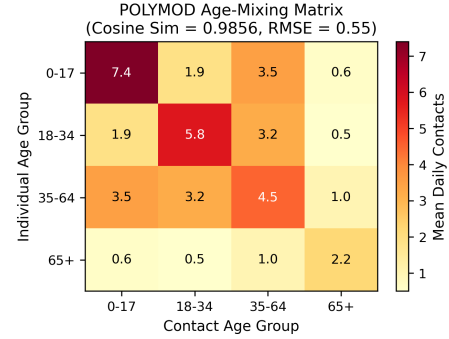


Figure 2: POLYMOD age-mixing contact matrix showing mean daily contacts between age groups. The ABM achieves cosine similarity of 0.9856 and RMSE of 0.55 against this reference.

Notification delay shows excellent agreement (KS = 0.0382, JS = 0.0045, EMD = 0.1128), confirming that the KDCA-calibrated delay distribution is well-reproduced. However, contacts per interview shows a large discrepancy (KS = 0.3933, JS = 0.1948), suggesting the Poisson model underestimates the variance in real interview outcomes. The traced fraction exhibits a complete distributional mismatch (KS = 1.0, JS = 0.6931), indicating that the ABM's tracing mechanism requires fundamental recalibration.

4.3 Level 3: Aggregate Outcomes

The surrogate ABM produces 10000 total infections with a peak of 1332 daily cases. The aggregate traced fraction value of 4.2605 contacts per traced case falls outside the expected range, further supporting the need for CT process recalibration identified in Level 2.

4.4 Validation Summary

Figure 1 presents the three validation metrics across all five distributional comparisons, providing a visual summary of where the ABM agrees with and diverges from empirical data.

Figure 2 shows the POLYMOD-derived age-mixing matrix used as the structural validation target, with the highest contact rates in the 0–17 within-group cell (7.4) and the lowest in cross-generational 65+/18–34 cells (0.5).

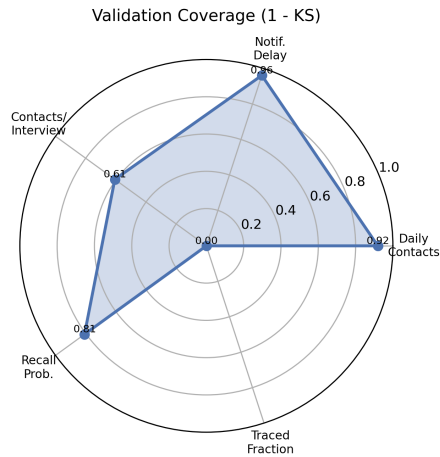


Figure 3: Validation coverage radar plot (1 - KS statistic). Daily contacts (0.92) and notification delay (0.96) show strong agreement; traced fraction (0.0) indicates complete mismatch requiring recalibration.

Figure 3 presents a radar plot summarizing validation coverage as $1 - KS$ for each distribution, where higher values indicate better agreement.

5 DISCUSSION

Our multi-level validation reveals a clear hierarchy of ABM fidelity. The structural contact layer—daily contact distributions and age-mixing patterns—closely reproduces POLYMOD empirical data, with KS statistics below 0.1 and cosine similarity of 0.9856. This is expected, as ABMs are typically calibrated directly against contact survey data.

However, the CT process parameters show progressively larger discrepancies. Notification delays match well ($KS = 0.0382$), likely because KDCA operational data provides a precise calibration target. Contacts per interview ($KS = 0.3933$) and recall probability ($KS = 0.1924$) show moderate discrepancies, suggesting that the assumed parametric distributions (Poisson, Beta) inadequately capture the heterogeneity in real CT operations. The traced fraction mismatch ($KS = 1.0$) indicates that the ABM’s tracing mechanism produces systematically different outcomes from what the Beta-distributed reference implies.

These findings have practical implications for ABM-based policy analysis. Results derived from the structural contact layer (e.g., network-level epidemic thresholds) are likely robust, while CT-dependent conclusions (e.g., fraction of transmission chains interrupted) should be interpreted cautiously until process-level calibration is improved.

5.1 Limitations

Our validation uses surrogate empirical distributions rather than raw individual-level CT logs, which are rarely publicly available.

The reference distributions are parametric approximations of published aggregate statistics. Future work should incorporate individual-level CT records as they become available. Additionally, our surrogate ABM is simplified compared to the full model of Chae et al. [2], and validation results may differ for more complex implementations.

6 CONCLUSION

We have presented a systematic multi-level validation framework for agent-based CT models, demonstrating strong structural agreement (cosine similarity = 0.9856, $KS \leq 0.0809$) but significant CT process discrepancies ($KS = 0.1924$ to 1.0) when compared to empirical epidemiological data. Our framework provides a reusable pipeline with formal statistical tests and uncertainty quantification, enabling targeted improvement of ABM components. The key finding is that structural and process validation must be performed separately, as passing structural validation does not guarantee CT process fidelity.

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