

Critical Challenges and Guidelines in Evaluating Synthetic Tabular Data: A Systematic Review

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Abstract. Generating synthetic tabular data can be challenging, however evaluation of their quality is just as challenging, if not more. This systematic review sheds light on the critical importance of rigorous evaluation of synthetic health data to ensure reliability, relevance, and their appropriate use. Based on screening of 1766 papers and a detailed review of 101 papers we identified key challenges, including lack of consensus on evaluation methods, improper use of evaluation metrics, limited input from domain experts, inadequate reporting of dataset characteristics, and limited reproducibility of results. In response, we provide several guidelines on the generation and evaluation of synthetic data, to allow the community to unlock and fully harness the transformative potential of synthetic data and accelerate innovation.

CCS Concepts: • Computing methodologies → Machine learning approaches; • Applied computing → Health informatics.

Additional Key Words and Phrases: synthetic data, tabular data, time series data

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1 INTRODUCTION

Access to high-quality data is fundamental to advancing scientific research. In disciplines such as healthcare, data is pivotal to enhance patient care, optimise resource management, and enable the discovery of new medical insights, particularly with the rise of artificial intelligence. Structured health data, such as tabular electronic health records, have been recognised as having one of the highest potential to provide timely and relevant information in clinical decision-making [1]. However, complex data-sharing governance rules have resulted in health data being locked away in isolated silos [2, 3], where they generally remain inaccessible except to a few researchers [4]. This inevitably hampers

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reproducible health research, hindering the advancement of patient care and impeding the future potential of clinical artificial intelligence [5].

There is an urgent need to democratise access to health data [4] without losing sight of patient privacy and confidentiality [6]. In this context, synthetic health data emerges as an attractive solution to address this challenge, and institutions worldwide are increasingly recognising their potential. For example, the United States Department of Health and Human Services has made available a Synthetic Health Data Generation Engine¹ to accelerate patient-centred outcomes research and “address the need for research-quality synthetic data”. The United Kingdom’s National Health Service (NHS) has rolled out an ‘Artificial Data Pilot’² that aims to “provide users with large volumes of data that share some of the characteristics of real data while protecting patient confidentiality”. Similar efforts are recorded in Canada’s health economic hub Health City [7] and Germany’s Charité Lab³ for Artificial Intelligence in Medicine. Research-quality synthetic data (the focus of our work) can be used to rapidly develop and test preliminary hypotheses before applying them to real datasets [8]. They can also improve research pipeline by acting as a proxy for real-world data [9]. Furthermore, the controlled generation of synthetic health data can include a balanced representation of different demographic groups [10]. This would ensure that the previously underrepresented socio-demographic groups are adequately represented, thereby mitigating biases in health research that arise from skewed real-world health datasets and, in turn, address model fairness [11–13].

However, despite the above-mentioned advantages of synthetic health data, major challenges remain with their large-scale adoption. One of the major challenge is the lack of consensus on evaluating synthetically generated data vis à vis the corresponding real data [14–16]. This not only makes it difficult to track the state-of-the-art progress of synthetic data generation methods but also poses barriers to trust and adoption, as well as presents regulatory and compliance issues.

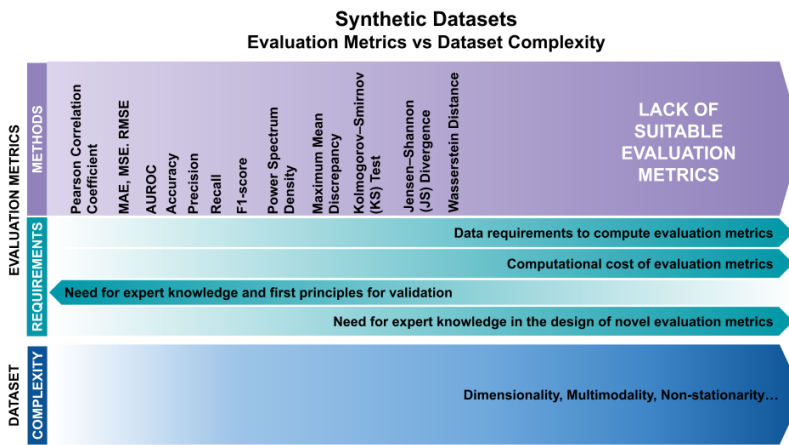


Fig. 1. There is an increasing lack of appropriate evaluation metrics (due to increasing difficulties in computation and increasing difficulties in evaluation of the metrics), with the increase in data complexity of the synthetic datasets.

To shed light on the evaluation approaches of synthetic data we have conducted a systematic review of 1766 research articles published in the last ten years.

This is the first review of this type and size to understand which approaches are being used to evaluate the quality of synthetic data, along with the associated data generation methods and their target application areas. We focused on structured tabular and time-series health data since this is one of the areas with the highest potential in advancing healthcare [17] and present

¹<https://aspe.hhs.gov/synthetic-health-data-generation-engine-accelerate-patient-centered-outcomes-research>

²<https://digital.nhs.uk/services/artificial-data>

³<https://claim.charite.de/en/>

unique data challenges, such as dealing with missingness. In addition, there is a higher consensus on the evaluation methods for other data modalities, such as imaging and text, where the respective research communities have developed metrics such as Fréchet Inception Distance (FID) [18] and BERTScore [19] respectively.

We observe that with the increasing complexity of the synthetic datasets (including dimensionality, multimodality, and non-stationarity), there is growing lack of suitable evaluation metrics, as shown in Fig.1. This is manifested in the increasing difficulties in the computation as well as evaluation of the metrics. Therefore, there's a critical need for the use of appropriate statistical evaluation metrics to critically evaluate complex synthetic data, including involvement of expert stakeholders in: i) selection of appropriate evaluation metrics and, ii) interpretation of the resulting outcomes. This type of collaboration between researchers and clinical practitioners can lead to development of methods and metrics that implicitly incorporate domain knowledge, resulting in decreased need for expert knowledge in evaluating future synthetic data. As a result, distilling domain knowledge into operational constraints and guaranteeing that the underlying medical processes that govern the data generation are safeguarded, will open the door to novel machine learning evaluation paradigms.

In the following section we show the results of our analysis, followed by guidelines in evaluating synthetic tabular data.

2 RESULTS

Based on the screening of 1766 papers and a detailed review of 101, we present the following results, grouped into four categories namely, evaluation, generation, purpose and impact of synthetic data, as well as reproducibility of the results.

2.1 Evaluation of Synthetic Data

We categorise the approaches used in the evaluation of synthetic data in: Direct vs Indirect approaches, and Quantitative vs Qualitative methods.

Direct evaluation approaches involve using existing, standardised metrics to assess the quality of the synthetic data. **Indirect evaluation approaches** include non-standardised, domain-specific methods (such as TSTR - Train on Synthetic, Test on Real) to assess synthetic data in real-world applications. Indirect approaches extend beyond the

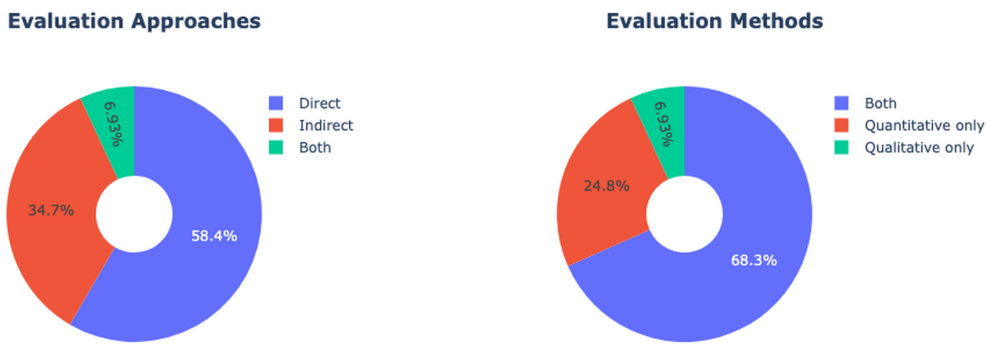


Fig. 2. (L) Breakdown of Evaluation Approaches into Direct vs Indirect Evaluation, and (R) Breakdown of Evaluation Methods into Quantitative vs Qualitative Methods

standard metrics and often include context-driven or subjective evaluations. We found that Direct approaches are the most common at 58.4%. About 34.7% of the publications use Indirect approaches. Additionally, 6.93% use both Direct and Indirect approaches in conjunction, to perform a holistic evaluation of their synthetic data (Fig. 2).

Quantitative evaluation methods give objective, measurable results and are crucial for ensuring that synthetic data aligns statistically with real data. They include **Statistical techniques** which use quantifiable metrics to compare synthetic data with the original data, and **ML-based techniques** which make use of classification and/or regression to assess how well synthetic data performs when used for specific downstream tasks. We found that Statistical evaluation is the most popular (54.5%), whereas ML-based techniques feature in 13.9% of all publications. About 24.8% publications use both Statistical and ML-based evaluation techniques in conjunction. (Fig. 3).

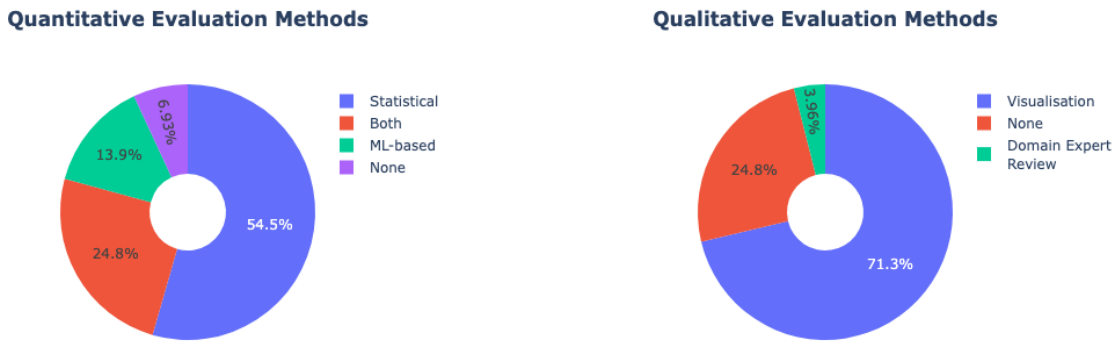


Fig. 3. (L) Breakdown of Quantitative Evaluation Methods into Statistical and ML-based methods, and (R) Breakdown of Qualitative Evaluation Methods into Visualisation and Domain Expert Review

The popularity of Statistical evaluation techniques remains high, however, a trend can be seen in publications using both Statistical and ML-based techniques together (Fig. 4).

Popularity Trend of Evaluation Methods

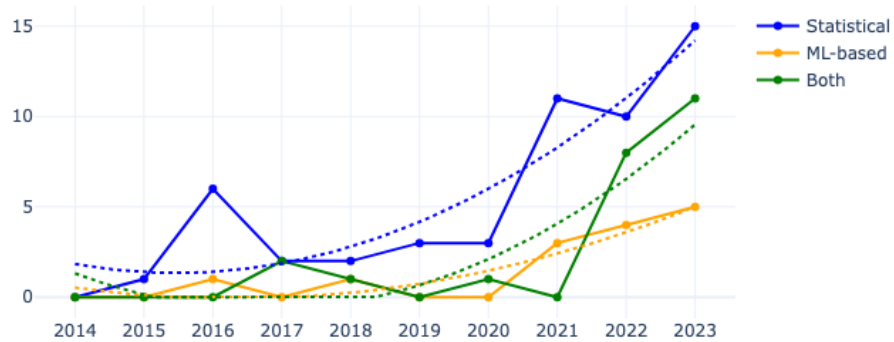


Fig. 4. Popularity trend of Statistical and ML-based Evaluation methods over the last decade, as obtained from publications included in this review. Dotted lines represent the overall polynomial trend.

The most used quantitative evaluation metrics include Jensen-Shannon (JS) distance, Pearson Correlation coefficient, and Maximum Mean Discrepancy (MMD), apart from the popular metrics such as AUC and F1-score for classification tasks, and Mean Square Error (MSE) and Root Mean Square Error (RMSE) for regression tasks (Fig. 5). We also note that the majority of the included papers (89.1%) use existing metrics, and only 10.9% of the publications use their own Author-defined metric for evaluation of the synthetic data.

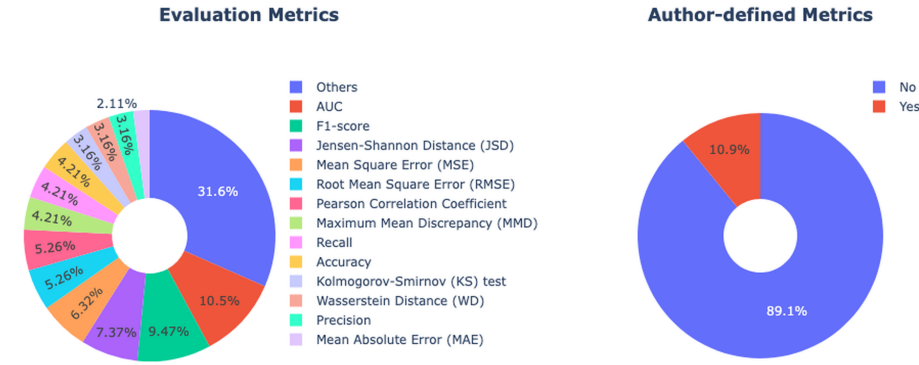


Fig. 5. (L) Most popular Evaluation Metrics, and (R) Breakdown of whether the metric is author-defined or not

Qualitative evaluation methods rely on subjective judgement and human interpretation to assess the quality of synthetic data. In the majority of the cases (71.3%), they take the form of **Visual Inspection** of the graphical representation of distributions of synthetic data (Fig. 3). We also note that despite its significance, use of **Domain Experts** as a qualitative evaluation method is not yet widely used and present only in a handful of all papers (3.96%).

Quantitative methods by themselves are used in about 24.8% of all publications, whereas for qualitative, this value is 5.93%. Most research (68.3%) uses a combination of both quantitative and qualitative methods. Fig. 6 gives a more detailed depiction of the most popular evaluation metrics and the papers utilising them that have been included in this review.

2.2 Generation of Synthetic Data

We categorise the models used for the generation of synthetic data into probabilistic and mechanistic models.

Probabilistic Models use statistical and probability distribution approaches to capture the statistical properties (such as distribution, correlations, and relationships between variables) of the real data, to generate the synthetic data. **Mechanistic Models**, on the other hand, use explicit rules, equations, or processes to simulate data based on how the underlying systems work. The most popular generation models are based on GANs, SMOTE, VAEs, Markov Chains, and Random Permutations (Fig. 7). Diffusion-based models are seeing a rise in popularity for longitudinal tabular data.

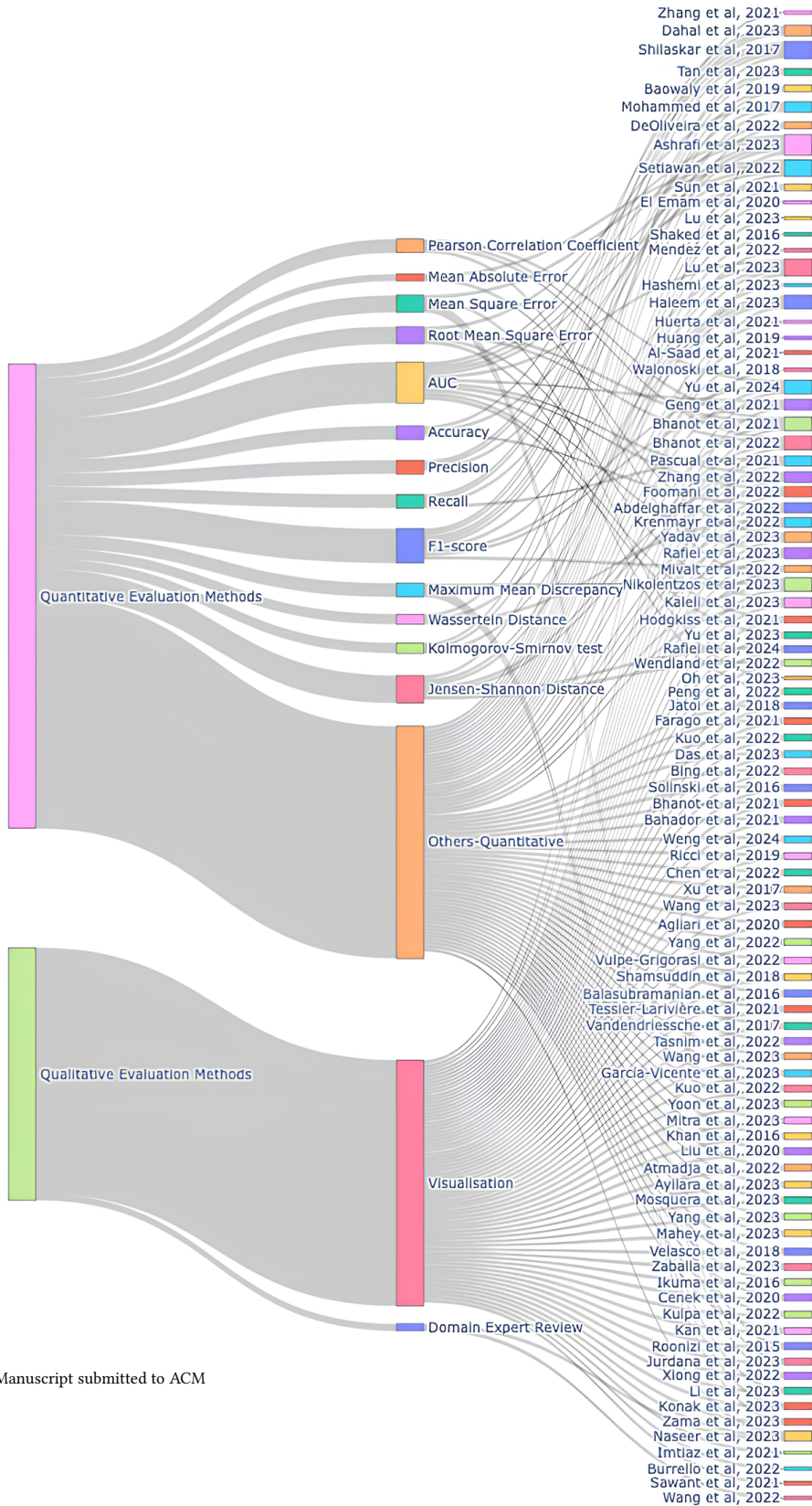


Fig. 6. Sankey depicting the most popular Evaluation Metrics and the papers utilising them, as obtained from the publications included in this review

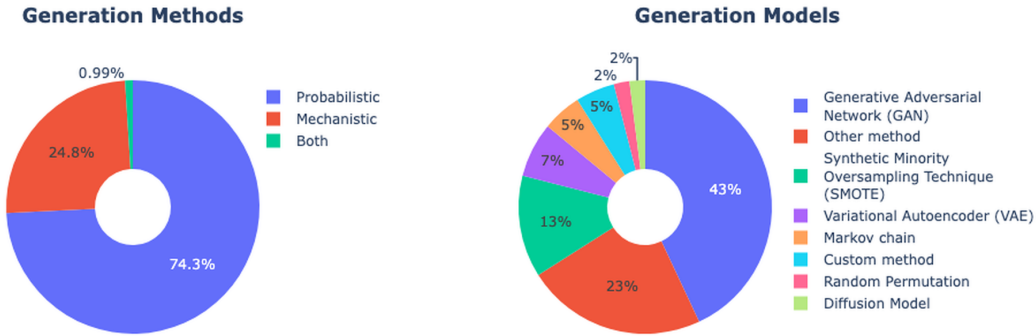


Fig. 7. (L) Breakdown of Generation Models into Probabilistic vs Mechanistic Models, and (R) Most popular Generation Models

We also observed a growing divergence between Probabilistic and Mechanistic models, with Probabilistic Models increasingly being more frequently used (74.3%) and Mechanistic Models tending to be more referenced in older publications only (Fig. 8).

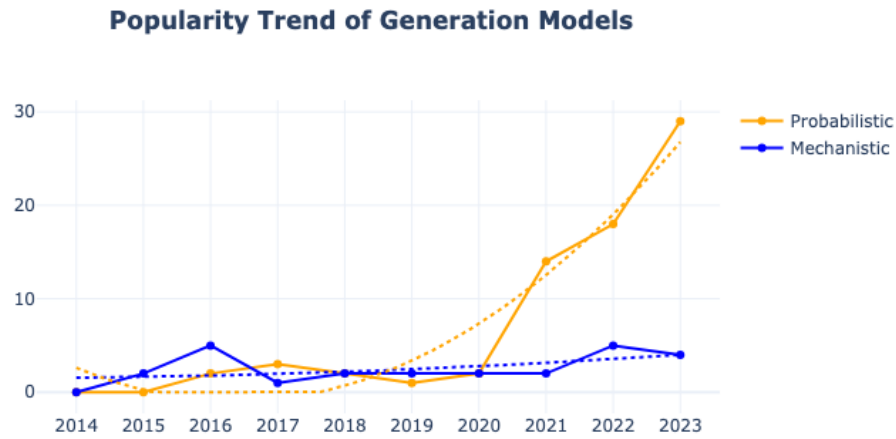


Fig. 8. Popularity trend of Probabilistic and Mechanistic Generation Models over the last decade, as obtained from publications included in this review. Dotted lines represent the overall polynomial trend.

2.3 Purpose and Impact of Synthetic Data

We found that privacy preservation, followed by predictive modelling and data-quality enhancement are the most popular objectives for the use of synthetic tabular data in healthcare. The most common diseases within the set of publications included in this review, for which synthetic health data is used, include those of the circulatory system, the nervous system, and neoplasms (Fig. 9). This may be driven by the popularity of the datasets, with MIMIC III[20] and MIMIC IV[21] being the most popular tabular and time-series health datasets.

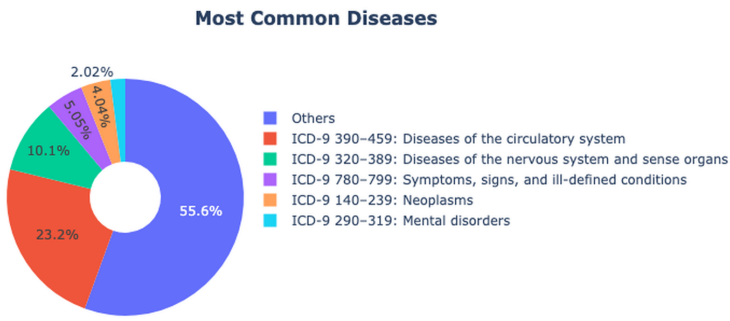


Fig. 9. Most common diseases (grouped by their ICD-9 codes), as seen through the publications included in this review.

search, the majority of the articles are published in journals and conferences with a primarily technical focus. Most of the research in synthetic health data is carried out in North America, followed by Europe and Asia, which may be influenced by the availability of health data and data protection regulations necessitating the use of synthetic data.

2.4 Reproducibility of Results

We define reproducibility as a factor of the use of publicly available datasets and the reporting of publicly accessible source code of the research. We note that, despite being a crucial piece of information, reproducibility is often not emphasised, and only 21.78% of all included publications were reproducible according to our definition. The majority of the papers (63.36%) use publicly available real-world health datasets, whereas about 12.87% used paid real-world health datasets for the creation of synthetic data. However, only 24.75% of the publications give details about their code along with a link to the code repository, which affects the overall reproducibility. Reproducibility is essential for ensuring the reliability and impact of scientific research. Particularly in healthcare where decisions can directly affect patient outcomes, reproducibility helps prevent errors, biases, and misleading conclusions. However, we found that it is an often overlooked aspect in most publications dealing with synthetic data.

In response to these results, we devise a set of reporting guidelines on the generation and evaluation of synthetic data, which are henceforth described in Section 3.

3 EVALUATION GUIDELINES FOR SYNTHETIC DATA

(1) Standardised Evaluation of Synthetic data: We found that synthetic data are sometimes used without a thorough assessment. When there is an assessment, we found that not only there is no consensus on the evaluation methods, but the chosen evaluation metrics are inconsistently applied. This makes an operational assessment of the entire process unreliable, thereby making it difficult to track state-of-the-art advancements and creates barriers to trust and the adoption of synthetic data.

For example, using Mean Square Error (MSE) metric as a measure of distortion to assess the validity of a synthetically generated waveform signal is appropriate but the validation needs to consider the particular

The widest-used repository is Physionet⁴ where these datasets are held, a testament to the value this repository provides to the research community. At the same time, this also poses a risk of perpetuating possible biases in the Physionet resources itself (for example, underrepresentation bias) to a global user base.

While there is some evidence to suggest that clinically oriented journals are also beginning to consider synthetic health data re-

⁴<https://physionet.org/>

features of the process. Two synthetic data generation methods with comparable MSE performances can still yield qualitatively very different signal features. Given the averaging that is implicitly performed in the computation of MSE, synthetic signals with uniformly distributed errors or distortions with respect to the real data will obtain similar MSE scores in comparison to signals with localised distortion patterns such as those modelled as impulsive noise. The signal characteristics in these two cases provide a stark contrast, and therefore, they will yield different conclusions in general when applied for validation or predictive tasks.

- (2) **Better Reporting of Dataset Characteristics:** Poor reporting of dataset details is a cause for concern since the type of data (such as categorical or continuous) and their distribution, significantly impact the quality of the generated synthetic data. Furthermore, potential biases may propagate in the synthetic data. We recommend improved reporting of dataset characteristics used for the generation of synthetic data.

For example, in developing a novel method, one would expect to assess its performance using real data or previously validated synthetic data. However, using synthetic data that has not been validated prior to the assessment of the proposed method does not provide robust evidence towards the validity of the method. Moreover, any claims about the validity of the synthetic data based on the performance of the proposed method are inherently inconclusive as they incur a circular reference problem that compromises the generated evidence.

- (3) **Prioritisation of Reproducibility of Results:** We emphasise the importance of clearly described evaluation metrics used, normalisation and aggregation of real data that were used as training datasets, and the entire experimental setup for the generation of synthetic data, including the chosen hyperparameters and the source code where possible. Reproducibility allows other researchers to validate and verify the claims of a study, and stakeholders, including clinicians and patients to develop trust in synthetic health data.

4 CONCLUSION

The potential of synthetic data to revolutionise Health AI research is immense, offering opportunities to address data scarcity, enhance privacy, and enable more robust model development. However, realising this potential requires a concerted effort to address critical challenges. Ensuring the applicability and fitness of synthetic data through rigorous assessment is paramount for its responsible use. Additionally, transparent reporting of datasets and ensuring reproducibility of results remain a cornerstone requirement of scientific progress.

Most importantly, the evaluation of synthetic health data must be guided by expert knowledge. As discussed earlier, domain expertise is critical for understanding the nuances of healthcare data and ensuring that synthetic datasets are technically sound, clinically relevant, and meaningful.

By adopting these guidelines and committing to ongoing collaboration, the health AI community can ensure that synthetic data is leveraged effectively and ethically, ultimately driving innovation and improving patient outcomes.

5 METHODOLOGY

In this section, we expand on our methodology for carrying out the systematic review. The Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) [22] statement provides a standardised framework for reporting systematic reviews. It consists of updated instructions on identifying, selecting, praising, and synthesising publications. Fig.10 outlines our methodology in accordance with the latest PRISMA guidelines.

5.1 Search Strategy

To establish an unambiguous search strategy, we laid out the following: (i.) The relevant databases used to search within a time frame, (ii.) The search terms to ensure comprehensive coverage of relevant studies, and, (iii.) The inclusion and exclusion criteria.

5.1.1 Databases and Search Engines. For this systematic review, we looked for publications on the following five databases: Scopus⁵, Web of Science⁶, PubMed⁷, IEEE Xplore⁸, and Association for Computing Machinery (ACM)⁹. Additional publications were also manually selected using Google Scholar.

5.1.2 Search Terms and Additional Limits.

Search Terms. We began by identifying publications dealing with synthetic data generation or augmentation. The focus was on finding publications that dealt with tabular or time-series data using wildcards in the search, and included all similar words including plurals and noun and verb forms of words. Keywords such as 'patient', 'health', and 'clinical' were also used to conduct the search within the health domain. As a result of the aforementioned considerations, the following search string was designed: (*synthe** OR *augment**) AND *generat** AND (*time-series* OR *time** OR *temporal**) AND (*tabular* OR *record**) AND (*patient** OR *medic** OR *health** OR *clinic** OR *ehr**), to be searched in the title-abstract-keywords or the topic field.

Additional Limits. We limited our search to publications in the last ten years, from January 1, 2014 to Jan 31, 2024. We also limited the search to peer-reviewed conferences and journal articles, written in the English language.

5.1.3 Inclusion and Exclusion Criteria.

Inclusion Criteria. The publications that were included in this systematic review met the following conditions:

- Publications that deal with tabular or time-series data.
- Publications that describe a method of generation of synthetic data and its evaluation against real data, or Publications that do not describe a method of generation of synthetic data but describe its evaluation vis a vis real data.
- Publications that deal with the generation of complete new synthetic datasets as well as the ones which deal with the augmentation of existing datasets with synthetic data.
- Peer-reviewed publications from journals and conferences. Strictly no pre-prints.

Exclusion Criteria. All possible publications that would be irrelevant to our study were excluded if they met any one of the following conditions:

- Publications that are not in the health domain.
- Publications that deal with image-, audio-, video-, or text-only modalities of data.
- Publications which themselves are narrative or systematic reviews.
- Publications on synthetic data that neither describe a method of generation of synthetic data nor its evaluation against real data.

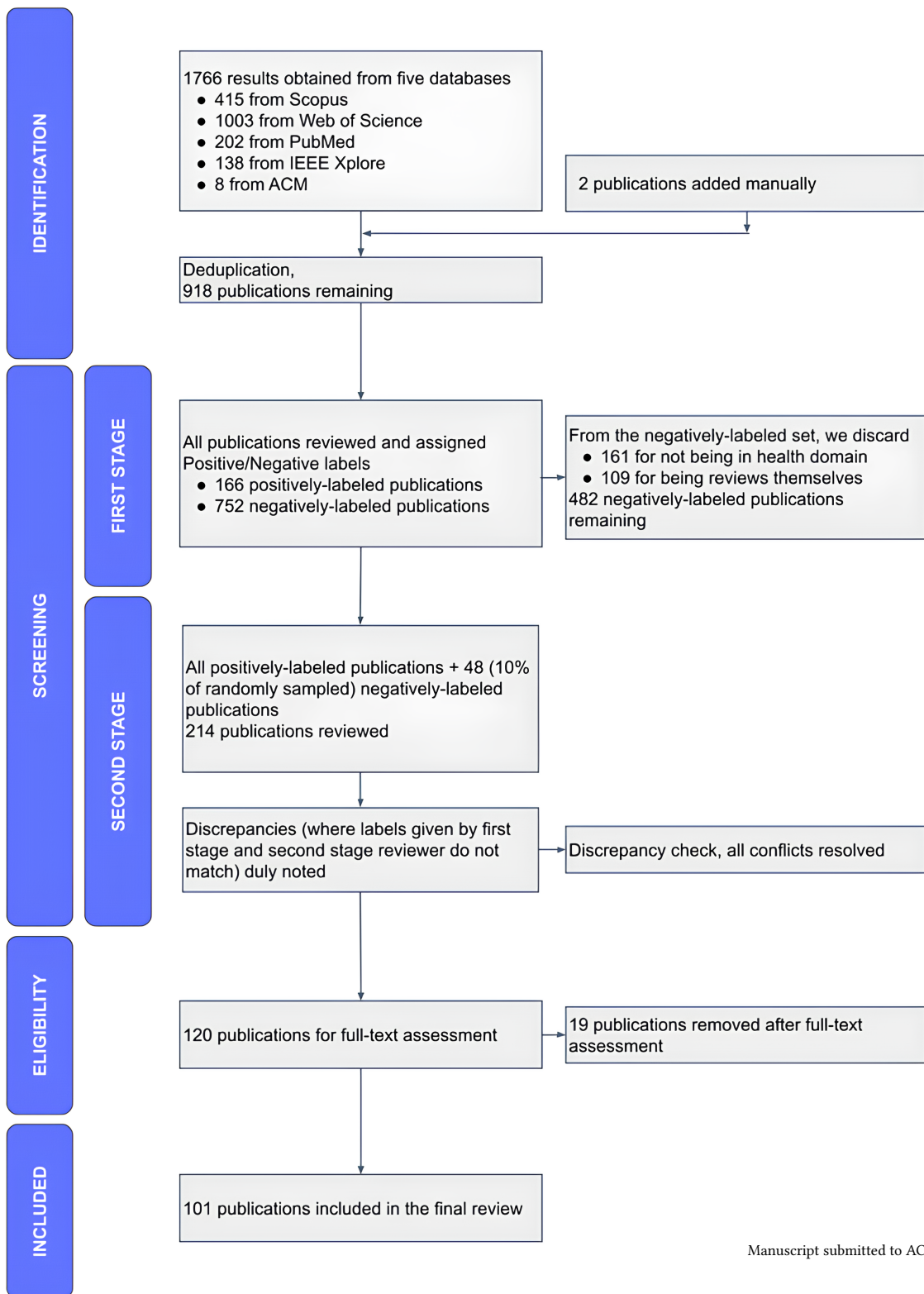
⁵<https://www.scopus.com>

⁶<https://www.webofscience.com>

⁷<https://pubmed.ncbi.nlm.nih.gov>

⁸<https://ieeexplore.ieee.org>

⁹<https://www.acm.org>



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Fig. 10. Popularity trend of synthetic tabular health data over the last decade, as obtained from publications included in this review

- Publications that describe a method of generation of synthetic data, but the synthetic data is not structurally similar (and therefore, not comparable) to real data.

5.2 Search Queries

Table 3 lists the search queries used across five databases respectively, and the number of relevant results obtained from each. Another set of limits on search engines were enforced via their graphical user interfaces (GUIs), hence they may have not been fully captured in the queries themselves. They are being mentioned here for the sake of completeness.

- The search was limited to publications in the last **ten years**, from January 1, 2014, to January 31, 2024.
- The search was limited to **peer-reviewed** conferences and journal articles written in the **English** language.

5.3 Selection Process

Based on the search strategy discussed in Section 5.1, 1766 publications were obtained from five search engines. This included 415 publications from Scopus, 1003 from the Web of Science, 202 from PubMed, 138 from IEEE Xplore, and 8 from ACM. Since many results were duplicated across databases, we carried out a deduplication process based on the DOIs of publications. As a result, 848 results were excluded and 918 publications remained for the first stage of assessment.

First Screening. The 918 publications obtained from search queries after deduplication, were divided among five reviewers with approximately 184 publications per reviewer. Each reviewer labelled the publications assigned to them Yes/No, signifying whether they thought the publication should be included or excluded along with the rationale for their decision. This gave us 166 publications labelled Yes, and 752 publications labelled No. From the 'No' set, we again discarded some publications, the most common reasons for their exclusion being that the research was not in the human health domain ($n=161$) or that the paper was a systematic review itself ($n=109$).

Next, we created a smaller subset of approximately 10% of the remaining No-labelled publications (482) and added it to all the Yes-labelled set. This combined set of 214 publications was used to perform a 'spot check' of labels: Publications were grouped by their first-stage reviewers and divided among the rest of the reviewers for a second round of reviewing. The first-stage reviewer's decision to include or exclude any particular publication was preserved but kept hidden from the view of the second-stage reviewers, to ensure that the reviewers' cognitive biases do not creep in during the labelling process, and that every publication gets assigned the correct label irrespective of who it was reviewed by in either of the two reviewing stages.

Second Screening. As with the first stage, each reviewer in the second stage provided a Yes or a No label to each publication in their set. No reviewer got to review the same publication in the second stage which they had already reviewed in the first stage. At the end of this exercise, any 'discrepancies' (cases where the labels given by the first-stage reviewer and the second-stage reviewer did not match) were duly noted. Then, a round of 'discrepancy checks' was carried out, where all reviewers looked at all discrepancies and provided feedback as to which of the two labels they agree with.

After a discussion on each occurrence of conflicting labels and resolving all discrepancies, we got the final labels for each publication. This resulted in a set of 120 publications for which the consensus of the reviewers was to include them in this systematic review.

Category	Data Items
Publication Details	DOI, Authors, Title, Publication Year, Journal/Conference Title
Synthetic Data Generation	Category of Generation Model (Mechanistic, Probabilistic), Name of Generation Method used, Purpose of Generation of synthetic data Disease/Disorder focused on, ICD-9 code
Synthetic Data Evaluation	Category of Quantitative Evaluation Method (ML-based, Statistical, Both, None), Category of Qualitative Evaluation Method (Visualisation, Others, None), Name of Evaluation Method used
Training Dataset Characteristics	Name, Size, Institution of Origin, Country of Origin, Visibility (Public/Private), Cost of Dataset Access
Source Code	Link to Source Code Repository

Table 1. Attributes against which data was collected for every publication

5.4 Data Items

For each of the 120 publications, an in-depth analysis was carried out. An additional 27 publications were excluded from the study upon full-text analysis, based on their relevance to this research.

Then, data was collected for the final 101 publications for 18 attributes which included: (i.) details about the publication including DOI, authors' names, title and year of publication and the details about the venue (journal/conference), (ii.) details specific to the generation of synthetic data such as the method (eg. WGAN-GP, Graph VAEs), its category (mechanistic/probabilistic), and purpose (eg. privacy preservation, clinical trial simulation), (iii.) details about the evaluation methods used, which includes the type of Quantitative evaluation (ML-based, Statistical or a combination), the type of Qualitative evaluation (for eg. Visualisation), the name of the method (eg. Jensen-Shannon divergence, Wasserstein distance) and the specific evaluation metrics used, (iv.) details about the dataset used in synthetic data generation, including the dataset name, size, institution and country of origin, and cost of dataset access, and (v.) details pertaining to the reproducibility of results including whether the dataset is openly accessible and if the source code has been made available.

A complete list of all data items against which data was captured is available in Table 1.

5.5 Reporting

The reporting of this systematic review adheres to the PRISMA guidelines [22]. We undertook measures to ensure transparency and reproducibility and facilitate critical appraisal and interpretation of the findings.

Risk of Bias Management: To establish the transparency of the findings and the results of this systematic review, we: (i.) used multiple databases to ensure no platform-specific bias creeps in, (ii.) used value-neutral search terms in the search query (iii.) got the publications reviewed by five reviewers in multiple screening stages (iv.) performed spot checks and discrepancy checks to ensure no reviewer-induced bias creeps in.

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A LIST OF PAPERS REVIEWED

An abridged version of the papers reviewed and their characteristics is provided in Table 2

Publication	Evaluation Metrics	Generation Methods	Dataset	Purpose
Jatoi et al, 2018 [23]	Custom method	Negative variational free energy, Localization error	Statistical Parametric Mapping - SPM12 software	Predictive modelling
Farago et al, 2021 [24]	Autoregressive modeling, Markov chain, RNN	Morphology, Mean, Variance, Autocorrelation, Power Spectral Density (PSD), Probability Distribution	Custom dataset	Signal quality analysis
Kuo et al, 2022 [25]	SAGAN	Accuracy, Standard Deviation	Custom dataset	Improve personalisation of prediction
Rafiei et al, 2023 [26]	CTGAN and SMOTE	AUC, AUROC, Sensitivity, Specificity, PPV, NPV, Bhattacharyya Distance	North Carolina Health System electronic medical record (EMR)	Fluid overload

DeOliveira et al, 2022 [27]	CTGAN and HARMGAN	Weighted Average F1-score, Ambiguity score	ExtraSensory dataset	Generating discrete synthetic data
Das et al, 2023 [28]	VAE	Dimension-Wise Probability, Bernoulli Success Probability, Counterfactual Digital Twin Evaluation, Presence Disclosure, Attribute Disclosure, Nearest Neighbor Adversarial Accuracy Risk	Phase III breast cancer clinical trial (NCT00174655), Small Cell Lung Carcinoma clinical trial dataset (NCT01439568),	Clinical trials
Bing et al, 2022 [29]	VAE	KNN	MIMIC-III	Mitigating representation bias
Bhanot et al, 2021 [30]	HealthGAN	Root Mean Square Error (RMSE), Pearson's Correlation Coefficient, Directional Symmetry, Short Time-Series Distance	American Time Use Survey (ATUS), Medical claims Autism Spectrum Disorder (ASD)	Privacy preservation, maintaining utility
El Emam et al, 2020 [31]	Conditional trees	Matching real with synthetic samples	Washington State Inpatient Database (SID) and Canadian COVID-19 case dataset	Privacy preservation
Lu et al, 2023 [32]	SMOTE	Precision, Recall, F1-score, Geometric mean, Area under the curve of the receiver operating characteristic curve (AUROC), Area under the precision-recall curve (AUPRC)	Taipei Medical University Hospital and Wan Fang Hospital (derivation), Taipei Medical University Shuang Ho Hospital (validation)	Predictive modelling
Solinski et al, 2016 [33]	Detrended fluctuation analysis (DFA)	Shannon Entropy, Poincare Plots, Multiscale Multifractal Analysis	Holter electrocardiogram (ECG) database and Complete electroencephalogram (EEG) recordings	NA
Bhanot et al, 2021 [34]	HealthGAN	Log Disparity, Time-Series Disparity	MIMIC-III and Average Sleep Time of Americans (ATUS)	Fairness
Shaked et al, 2016 [35]	Markovian Model	Mean Similarity, Intersection	MIMIC-III	Privacy preservation
Bahador et al, 2021 [36]	DE-NLPCA	Accuracy	Activities of daily living (ADL) dataset, and EEG / ECG dataset from Northern Ostrobothnia Hospital	Predictive modelling
Weng et al, 2024 [37]	MVIIIL-GAN	Missing Values Reconstruction Error	MIMIC-IV	Dataset balancing
Bhanot et al, 2022[38]	Bootstrapping, Random Permutation and HealthGAN	Root Mean Square Error (RMSE), Pearson's Correlation Coefficient, Short Time-Series Distance (STS), Directional Symmetry (DS)	American Time Use Survey (ATUS) dataset and Autism Spectrum Disorder (ASD) claims dataset.	Addressing Data unavailability, Privacy preservation

Nikolentzos et al, 2023 [39]	Variational Graph Autoencoder (VGAE)	Weisfeiler-Lehman (WL) Graph Kernel, Shortest Path (SP) Graph Kernel, Graph Kernel-Maximum Mean Discrepancy (GK-MMD), Pearson Correlation Coefficient	Subtree MIMIC-IV	Privacy preservation
Ricci et al, 2019 [40]	Custom method	Poisson Sequence, Correlated Heartbeatlike Sequence, Hénon Map Sequence	Electronic oscillator sequence, Heartbeat sequence, and Neural sequence	NA
Hodgkiss et al, 2021 [41]	Custom method	AUC	Normal Sinus Rhythm Dataset	Cybersecurity
Chen et al, 2022 [42]	CTGAN	Classifier	CHB-MIT EEG dataset	Dataset balancing
Xu et al, 2017 [43]	Custom method	Markov Chain, Vector Auto-Regressive Model, Continuous-Time Markov Chain, Logistic Regression, Hawkes Processes, Modulated Poisson Processes, Self-Correcting Process	MIMIC-II	Dataset balancing
Mohammed et al, 2017 [44]	ARMA	F1-score, AUC, NOP	Custom dataset	Addressing Data unavailability
Wang et al, 2023 [45]	AFE-GAN (Atrial Fibrillation-like ECG GAN)	Two of the four winning atrial fibrillation detectors from the 2017 PhysioNet Challenge - Hong detector, Datta detector	training set from the 2017 PhysioNet Challenge	Addressing Data unavailability
Imtiaz et al, 2021 [46]	BGAN (Boundary-seeking GAN)	Visualisation only	Custom dataset - from Fitbit Charge 2 HR smartwatches	Privacy preservation
Ashrafi et al, 2023 [47]	simpleGAN, medGAN, DopelGANer, DPGAN, and PPGAN	F1-score, Precision, Recall, Root Mean Square Error (RMSE), AUC, Attacker Advantage	(Patient interactions with a tablet game (PflegeTab))	Privacy preservation
Rafiei et al, 2024 [48]	SMOTE, CTGAN	Jensen-Shannon Divergence (JSD), Bhattacharyya Distance, Mann-Whitney U Test, Benjamini-Hochberg (BH) procedure	Custom dataset	Predictive modelling
Agliari et al, 2020 [49]	Custom method	Power Spectrum Density (PSD)	Custom dataset	Method evaluation

Li et al, 2023 [50]	EHR-M-GAN and EHR-M-GANconditional	Maximum Mean Discrepancy (MMD), Dimension-Wise Probability, Discriminative Score, Patient Trajectories, Pearson pairwise correlations, Autocorrelation function, Membership Inference Attack, Differential Privacy	MIMIC-III, eICU and HiRID	Addressing Data unavailability, Privacy preservation
Zhang et al, 2022 [51]	LS-EHR	Jensen–Shannon Divergence (JSD), AUC	Custom datasets (two)	NA
Oh et al, 2023 [52]	Monte Carlo simulations	Relative Bias, Confidence Limit Ratios (CLRs), Mean Square Error (MSE)	Custom dataset (South Korea's patients' healthcare resource utilization database)	Checking bias
Burrello et al, 2022 [53]	DA techniques and DL HR algorithms	Visualisation only	PPGDalia	Health monitoring
Mendez et al, 2022 [54]	GAN	Welch's Test	NA	Health monitoring and Data privacy
Yang et al, 2022 [55]	Physics-based models	DFA	PhysioNet	Predictive modelling
Vulpe-Grigorasi et al, 2022 [56]	GAN	RMSD, SDNN	PhysioNet	Increased diagnosis accuracy
Shamsuddin et al, 2018 [57]	Virtual Patient Model	NB, SVM and TB	ARem and EEG	Addressing Data unavailability
Balasubramanian et al, 2016 [58]	MDMs (Multidimensional Motifs)	Graph Clustering Method	Electromagnetic Articulography and Motion Capture and Muscle Activity	Personalised diagnosis and therapy
Tessier-Larivière et al, 2021 [59]	PNS-GAN	Power Spectral Density, Euclidean distance	BIOS-IT3 Dataset	Data Augmentation
Vandendriessche et al, 2017 [60]	MSE	Classifier	MIMIC-III (heart and sepsis)	Predictive modelling
Tasnim et al, 2022 [61]	SMOTE	Classifier	BCIAUT-P300	Addressing Health inequality
Wang et al, 2023 [62]	SMOTE and WCGAN-GP	Classifier	ImmPort (Immunology Database and Analysis Portal) data	Enhancing health clinical data
García-Vicente et al, 2023 [63]	SMOTE, CTGAN, TVAE	LASSO, SVM, KNN, DT	Norwegian Centre for E-health Research	Data quality enhancement
Kuo et al, 2022 [64]	GAN	Classifier	MIMIC-III and EuResist23	Data quality enhancement and Privacy preservation

Yoon et al, 2023 [65]	Sequential encoder-decoder methods and GAN	Classifier	MIMIC-III	Privacy preservation
Mitra et al, 2023 [66]	CardioSim PC-based system	Classifier	mitdb	Data quality enhancement
Khan et al, 2016 [67]	NA	MEWMA	accelerometer	Health and well-being assessment
Roonizi et al, 2015 [68]	SHVR	Mean Square Error (MSE), Wilcoxon Rank Test	synthetic ECG	Predictive modelling
Konak et al, 2023 [69]	TimeGAN and Animations	MMD	PAMAP2 and SONAR-LAB	Addressing Data unavailability
Tan et al, 2023 [70]	TabGAN and SMOTE	Jensen–Shannon Divergence (JSD), Wasserstein Distance (WD), Diff Corr	SUPPORT and METABRIC	Predictive modelling
Liu et al, 2020 [71]	Deep Sequential Weighting (DSW)	LR, RF, KNN, PSM, CFR, CF, BART	Custom dataset and MIMIC-III	Predictive modelling
Atmadja et al, 2022 [72]	GAN	CNN	MIT-BIH	Predictive modelling
Hashemi et al, 2023 [73]	GAN	PCA, t-SNE, pairwise correlations, RNN	Sins, MIMIC-VI	Privacy preservation
Wang et al, 2022 [74]	Markov Jump Process	Domain Expert Review	NA	Predictive modelling
Huerta et al, 2021 [75]	Standard Data Augmentation Transformations	McNemar test	PhysioNet/CinC Challenge 2017 database	Addressing Data unavailability
Ayilara et al, 2023 [76]	OSIM2 and ModOSIM	Concordance Correlation Coefficient	Population Research Data Repository (PRDR)	Method evaluation
Haleem et al, 2023 [77]	TC-Multi GAN and Document Sequence Generator	Wasserstein Distance, Kolmogorov-Smirnov Test, Jensen-Shannon Distance, Distance Pairwise Correlation, Sample Kernel Density Estimations	GATEKEEPER EU project	Addressing Synthetic Data feasibility
Zama et al, 2023 [78]	Diffusion-based model	Dynamic Time Warping, Maximum Mean Discrepancy	PTB-XL	Privacy preservation
Mosquera et al, 2023 [79]	RNN with LTSM and GRU	Hellinger's Distance, Cox Regression Hazard Ratios	Alberta Health's administrative dataset	Addressing Synthetic Data feasibility
Huang et al, 2019 [80]	Delete, update, switch operations	Pairwise Similarity Score	Rochester epidemiology project	Predictive modelling
Dahal et al, 2023 [81]	EC-WCGAN	Precision, Recall, F1-score	AHADB, VFDB and CUDB	Dataset balancing
Setiawan et al, 2022 [82]	SMOTE	Accuracy, Sensitivity, Specificity, ROC, Cross-Validation, MSE, MAE	PhysioNet Apnea-ECG data-base (PAED)	Dataset balancing

Jurdana et al, 2023 [83]	Custom method	MSE	EEG, Royal Brisbane	Predictive modelling
Yang et al, 2023	TS-GAN	LSTM-based Discriminator, Discriminator loss, Maximum Mean Discrepancy, Principal Component Analysis, t-SNE, Sequence diagrams, Accuracy	ECG_200, NonInvasiveFetalECG_Thorax1, and mHealth	Data augmentation
Mahey et al, 2023 [84]	Simulation	Channel by Channel Covariance, EOG, 1/f Function, Spatial Covariance	NA	Addressing Data unavailability
Velasco et al, 2018 [85]	Evolutionary algorithm	Wilcoxon Rank Sum Test (Mann Whitney Wilcoxon) (MWW)	Principe de Asturias Hospital	Addressing Data unavailability
Xiong et al, 2022 [86]	Custom method	Mean Square Error (MSE), Average Standard Deviation, Frequency Distribution	PhysioNet 2017 challenge dataset	Dataset balancing
Krenmayr et al, 2022 [87]	GAN with bi-LSTM	Euclidean Distance, Wasserstein Distance	NA	Addressing Data unavailability
Zaballa et al, 2023 [88]	Probabilistic generative model (HMM and EM)	Average Log Likelihood	NA	Predictive modelling
Wendland et al, 2022 [89]	Multimodal Neural Ordinary Differential Equations	Jensen-Shannon Divergence	PPMI (Parkinson), and NACC (Alzheimer)	Predictive modelling
Kaleli et al, 2023 [90]	GAN with CNN and Transformer	Percent Root Mean Square Difference (PRD), Root Mean Square Error (RMSE), Frechet Distance (FD)	MIT-BIH dataset	Privacy preservation, Predictive modelling
Sun et al, 2021 [91]	Longitudinal GAN	AUROC, AUPCR, AUC	Cerner Health Facts database	Predictive modelling, Privacy Preservation
Shilaskar et al, 2017 [92]	Resampling, modified Particle Swarm Optimization	Accuracy, Precision, Recall, Sensitivity, F1-score	Vani Dataset, Thyroid Dataset, PdA, Cleveland, Audiology, SVD, Vertigo	Predictive modelling
Foomani et al, 2022 [93]	GAN	Jensen-Shannon Divergence, AUC	EMR data from Vascular Centers, Milwaukee, WI	Predictive modelling
Peng et al, 2022 [94]	Gaussian Kernels	Root Mean Square Error (RMSE)	NA	Predictive modelling
Geng et al, 2021 [95]	GAN	AUC, F1-score	NA	Predictive modelling
Mivalt et al, 2022 [96]	GAN	Cohen's Kappa, F1-score	Multicenter Intracranial EEG Dataset	NA
Pascual et al, 2021 [97]	GAN	Cosine Similarity, Recall	EPILEPSIAE	Privacy preservation

Lu et al, 2023	GAN [98]	Jensen-Shannon Divergence, Normalised Distance, AUC, F1-score	MIMIC-III and MIMIC-IV	Privacy preservation
Ikuma et al, 2016 [99]	Karhunen-Loeve transformation, Time-series model perturbations	Correlation analysis relative vibration power represented by a synthetic waveform	NA	Predictive modelling
Zhang et al, 2021 [100]	GAN	Diagnosis forecast analysis, Kolmogorov-Smirnov Test	Synthetic Derivative at Vanderbilt University Medical Center	Privacy preservation
Baowaly et al, 2019 [101]	GAN	Dimension-Wise Average, Kolmogorov-Smirnov (KS) Test, Association Rule Mining	MIMIC-III and NHIRD	Predictive modelling
Abdelghaffar et al, 2022 [102]	GAN	Relative Entropy, Accuracy	Wadsworth BCI Dataset from the BCI competition III	Predictive modelling
Al-Saad et al, 2021 [103]	DPGAN	Dimension-Wise Average, AU-ROC, Area under the Precision-Recall Curve, Accuracy	Arizona State's Kinesiology Department	Predictive modelling, Privacy preservation
Yu et al, 2023 [104]	GAN	AUC	CHB-MIT dataset	Predictive modelling
Cenek et al, 2020 [105]	Frequency Domain Model	NA	CIRCADA-S	Predictive modelling
Yadav et al, 2023 [106]	GAN	Mean Absolute Error (MAE), MRLE, PCA, t-SNE	UNIMIB	Predictive modelling
Yu et al, 2024 [107]	Temporal Convolutional Network	Dipole Localization Error (DLE), Normalized Hamming Distance, Sensitivity, Specificity, False Detection Rate, F1-score, Pearson Correlation	NA	Predictive modelling
Walonuski et al, 2018 [108]	Markovian model (PADARSER)	Prevalence Difference Error	Multiple	Predictive modelling
Kulpa et al, 2022 [109]	Autoregressive model, Markov chain, RNN	Power Spectral Density (PSD)	MIT-BIH NSTDB	Predictive modelling
Kan et al, 2021 [110]	GAN	Average Error Rate	Temple University Hospital Abnormal EEG Corpus	Predictive modelling
Naseer et al, 2023 [111]	Continuous-Time Diffusion Models	Dimension-wise distribution, Pairwise Correlation difference, Log-cluster, Synthetic ranking agreement, Membership Inference Attack, Blinded Clinician Evaluation, Domain Expert Review	MIMIC-III and ED-EHR datasets	NA

Qian et al, 2024 [112]	DPGAN, PATEGAN), ADSGAN	Fidelity (Alpha-Precision), Diversity (Beta-Recall), Authenticity, Wasserstein distance, Jensen-Shannon distance, Inverse Kullback-Leibler divergence, Chi-Squared Test, Kolmogorov-Smirnov test, k-anonymity, DOMIAS AUC	Ever-smokers in UK Biobank Database	Privacy preservation
Lu et al, 2023 [32]	SMOTE	Decision Tree, Random Forest, Logistic Regression, Extreme Gradient Boosting, Support Vector Machines	Custom data	Dataset balancing
Sawant et al, 2021 [113]	SMOTE	Sensitivity, Specificity, Overall score (Average of Sensitivity and Specificity)	PhysioNet / CinC challenge 2016, and PASCAL	Dataset balancing
Akter et al, 2021 [114]	SMOTENC (SMOTE variant)	Classifier	Quantitative Checklist for Autism in Toddlers-10 (CHAT-10), and Autism Spectrum Quotient-10 (AQ-10)	Predictive modelling
Katekarn et al, 2023 [115]	Custom method	Participant satisfaction questionnaire and SPSS	Custom dataset	Method evaluation
Vemuri et al, 2016 [116]	Custom method	Custom evaluation metrics (Measurement of Uncertainty, Measuring Uncertainty in Endoscope Tip)	NA	Predictive modelling
Valdano et al, 2015 [117]	Custom method	Visualisation only	NA	Disease modelling
Covioli et al, 2023 [118]	Simulation	Visualisation only	MIMIC-III and MIMIC-III waveform matched dataset	Method evaluation
Tomek et al, 2016 [119]	Cellular automata	Median, p-value	Custom dataset	Method evaluation
Squires et al, 2022 [120]	Python packages (random and fake)	Custom evaluation metrics	Custom dataset	Addressing Data unavailability
Oliveira et al, 2023 [121]	CGAN and CT-GAN	False Positives, False Negatives, Accuracy, Specificity, Sensitivity, AUC	PARK Facial Mimic	Data Augmentation

B SEARCH QUERIES

The following is a list of search queries used across five databases, and the number of relevant results obtained. It should be noted that additional filtering criteria were set on these databases, including the date of publication range (2014-2024) and the language of the publication (English).

Database	Query	#
Scopus	(synthe* OR augment*) AND generat* AND (time-series OR time* OR temporal*) AND (tabular OR record*) AND (patient* OR medic* OR health* OR clinic* OR ehr*)} on title-abstract-keywords	415
Web of Science	(synthe* OR augment*) AND generat* AND (time-series OR time* OR temporal*) AND (tabular OR record*) AND (patient* OR medic* OR health* OR clinic* OR ehr*)} (Topic)	1003
PubMed	(synthe* [Title/Abstract] OR augment* [Title/Abstract]) AND generat*[Title/Abstract] AND (time-series[Title/Abstract] OR time*[Title/Abstract] OR temporal*[Title/Abstract]) AND (tabular[Title/Abstract] OR record* [Title/Abstract]) AND (patient*[Title/Abstract] OR medic*[Title/Abstract] OR health*[Title/Abstract] OR clinic*[Title/Abstract]) on title-abstract]	202
IEEE Xplore	("All Metadata":synthe* OR "All Metadata":augment*) AND ("All Metadata":time* OR "All Metadata":temporal*) AND ("All Metadata":tabular OR "All Metadata":record*) AND ("All Metadata":patient* OR "All Metadata":medic* OR "All Metadata":health* OR "All Metadata":clinic* OR "All Metadata":ehr*) AND ("All Metadata":generat*)	138
ACM	[Abstract: synthe*] AND [[Abstract: time*] OR [Abstract: temporal*]] AND [[Abstract: patient*] OR [Abstract: medic*] OR [Abstract: health*] OR [Abstract: clinic*] OR [Abstract: ehr*]] AND [E-Publication Date: (01/01/2014 TO 31/01/2024)]	8

Table 3. Search Queries used, and the number of Results obtained from each Database