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RESEARCH ARTICLE

CAN DIGITAL NEUROPHENOTYPING AND ARTIFICIAL INTELLIGENCE TRANSFORM THE EVALUATION OF CRANIAL OSTEOPATHIC INTERVENTIONS

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Abstract

Cranial osteopathic interventions occupy a contested space in manual medicine, with longstanding debates centering on the reproducibility of proposed mechanisms, the validity of inter-rater reliability in the detection of craniosacral rhythms, and the absence of objective physiological outcome measures. Despite widespread clinical application, scientific evaluation has been impeded by reliance on subjective self-report and palpatory endpoints. Emerging methodologies in digital neurophenotyping, encompassing multimodal wearable biosensors, continuous electroencephalography, heart rate variability analysis, digital cognitive assessment, and actigraphy-based sleep monitoring, now offer the potential to capture dynamic neurophysiological states with previously unattainable precision and ecological validity. When integrated with machine learning frameworks capable of processing high-dimensional, temporally rich datasets, these technologies may provide the objective biomarker infrastructure necessary to design scientifically credible trials of cranial osteopathic practice. This commentary examines the conceptual and methodological basis for such an approach, proposes a translational research roadmap, and identifies priority challenges, including signal interpretability, ethical governance, and the risk of technological overreach. The intention is not to advocate for cranial osteopathy's efficacy but to argue that the measurement gap represents the most tractable problem for the interdisciplinary research community to address.

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Introduction:-

Cranial osteopathy, also termed craniosacral osteopathy or cranial technique within the osteopathic curriculum, encompasses a family of low-force manual interventions applied to the skull, spinal column, and sacrum.[1] First codified by William Garner Sutherland in the early twentieth century, the approach rests on the hypothesis that rhythmic, subtle movements of the cranial bones and cerebrospinal fluid express a palpable primary respiratory mechanism whose normalisation may yield therapeutic benefit.[1] Despite widespread clinical use across osteopathic and physiotherapy settings in Europe, Australia, and increasingly South Asia, the discipline occupies a contested position between clinical tradition and scientific scepticism.[2,1] Systematic reviews have documented insufficient evidence of efficacy for most claimed indications, while proponents argue that adequate measurement instruments have not yet been applied.[2]

The crux of the problem is methodological. Clinical assessment of cranial osteopathic interventions has historically relied on practitioner palpation, patient-reported outcome measures, and observational endpoints, tools inherently vulnerable to detection bias, expectation effects, and regression to the mean.[2,1] Where objective physiological recording has been attempted, study designs have been underpowered, and instrumentation has been insufficiently sensitive to detect the subtle, distributed neurophysiological changes that proponents propose.[2]

Two converging technological developments now create a genuine opportunity to revisit this measurement gap. Digital neurophenotyping, the systematic, longitudinal characterisation of neurological function through passively and actively acquired digital data streams, has matured rapidly, propelled by advances in wearable sensor engineering, mobile health platforms, and open-source signal processing pipelines.[3,4,5] In parallel, machine learning methods capable of extracting structured patterns from multimodal physiological time-series have achieved validated utility in neurological disease stratification, treatment response prediction, and biomarker discovery.[9,12] Together, these capabilities may finally provide the scientific infrastructure required to ask precise, falsifiable questions about what, if anything, changes in neurophysiology following cranial osteopathic intervention.[3,11]

Why Objective Measurement Matters:-

The history of medicine is in large part a history of measurement. The introduction of sphygmomanometry transformed hypertension from a symptomatic cluster into a quantifiable risk factor amenable to titrated pharmacotherapy.[7] Neurological conditions once classified by descriptive phenomenology were transformed by dopamine transporter imaging and accelerometry into disorders with objective, stratifiable biomarkers.[9] In each case, objective measurement did not merely confirm clinical observation; it revealed heterogeneity, stratified populations, refined mechanism hypotheses, and enabled dose-response characterisation that had previously been inaccessible.[7,9,2]

The problem of subjective endpoints in manual medicine is well-characterised. Inter-rater reliability studies of craniosacral rhythm detection have produced consistently poor kappa values, with some investigations reporting agreement no better than chance.[1] Patient-reported outcome measures capture the patient experience authentically but cannot distinguish specific from non-specific effects, cannot be adequately blinded in active-comparator designs, and are insufficient to support mechanistic inference.[2] These limitations do not imply that clinical benefit does not occur; they imply that the evidentiary architecture required to detect and characterise benefit is absent.[2,1]

The parallel with early psychiatric and neurodevelopmental research is instructive. For decades, the absence of objective biomarkers for conditions such as major depressive disorder and post-traumatic stress disorder constrained trial design to symptom rating scales, creating a literature systematically underpowered to detect neurobiological heterogeneity.[3] The emergence of neuroimaging, polysomnography, and actigraphy as supplementary outcome measures substantially elevated the precision and scientific credibility of research in these fields.[8,9] Manual medicine currently occupies an analogous position and may benefit from analogous solutions.[10,2]

Digital Neurophenotyping: A New Opportunity:-

Digital neurophenotyping refers to the use of digitally acquired behavioural, physiological, and cognitive data, collected continuously or episodically in naturalistic settings, to characterise neurological and neuropsychiatric states with temporal and ecological fidelity unavailable in single-session laboratory assessments.[3,4,5] The term was formalised in the neuropsychiatric context by Insel and Cuthbert, who argued that precision medicine required the systematic capture of digital phenotypic markers beyond clinical diagnosis.[3] It has since expanded to encompass wearable-derived cardiac, kinematic, sleep, and cognitive data streams across a spectrum of neurological and systemic conditions.[4,5]

Electroencephalography (EEG) provides perhaps the most direct window into cortical activity relevant to cranial osteopathic hypotheses. Advances in dry-electrode, head-mounted EEG systems now permit ambulatory recording with sufficient signal quality for spectral and connectivity analysis.[9] Power spectral density across delta, theta, alpha, and beta bands, together with measures of functional connectivity such as coherence and phase-amplitude coupling, are established surrogates of cortical arousal state and have demonstrated sensitivity to a range of non-pharmacological interventions.[9] Whether cranial osteopathic treatment produces measurable, reproducible shifts in these indices is an empirically open question, but it is now an answerable one given available instrumentation.[9,1]

Heart rate variability (HRV) analysis offers a non-invasive index of autonomic nervous system tone extensively validated as a biomarker of stress reactivity, vagal efference, and allostatic load.[6] High-frequency HRV power, reflecting parasympathetic modulation of sinoatrial node firing, is sensitive to relaxation interventions, manual therapy, and biofeedback in controlled settings.[6] Consumer-grade photoplethysmography devices now deliver HRV metrics of research-grade validity in free-living conditions, enabling before-after-follow-up designs with ecological validity that laboratory sessions cannot replicate.[6,7]

Sleep architecture, quantified through actigraphy or consumer polysomnography, represents a systems-level integrator of autonomic, neuroendocrine, and restorative processes.[8] Similarly, respiratory pattern analysis, including respiratory rate variability and nocturnal breathing metrics, may capture changes in brainstem or diaphragmatic regulation plausibly linked to cranial osteopathic mechanisms.[1] Validated digital cognitive assessment platforms, deployable via smartphone in under ten minutes, provide reliable measures of processing speed, working memory, sustained attention, and executive function that are sensitive to autonomic state and fatigue.[4,5]

The Role of Artificial Intelligence:-

The individual data streams described above generate physiological time-series of considerable dimensionality, temporal complexity, and inter-individual variability. Classical univariate statistical approaches, comparing group means on a single outcome at a single time point, are poorly suited to this data structure and have contributed to replication failures that characterise much of the existing manual medicine literature.[2] Machine learning frameworks offer complementary analytical capabilities that align more naturally with the heterogeneous, multivariate, temporally extended data that digital neurophenotyping generates.[11,12]

Supervised learning methods, including regularised regression, support vector machines, and gradient boosted trees, have demonstrated utility in predicting treatment response from pre-treatment physiological profiles in psychiatric and rehabilitation contexts, opening the possibility of responder-stratification that could substantially increase statistical power in cranial osteopathy trials through enriched enrolment designs.[11,12] Unsupervised clustering and manifold learning methods, such as UMAP and variational autoencoders, offer data-driven identification of physiological subtypes among participants without imposing a priori categories, which is well-suited to a field where the relevant population heterogeneity is poorly characterised.[11]

Temporal modelling methods, including recurrent neural networks, long short-term memory architectures, and transformer-based models for physiological time-series, have achieved state-of-the-art performance in automated detection of sleep staging, seizure prediction, and autonomic state classification.[11,12] Applied to pre- and post-intervention neurophenotyping data, such architectures could identify trajectory patterns, including onset timing, duration, and magnitude of change, that simple before-after comparisons cannot resolve.[11]

It is essential, however, to enumerate the limitations of AI-assisted analysis with equal candour. Contemporary deep learning models are vulnerable to overfitting in small clinical samples, a near-universal characteristic of early-phase manual medicine trials.[11] Interpretability remains a contested challenge: a model that accurately classifies responders does not necessarily identify the mechanistic pathway responsible for classification, and clinical inference drawn from opaque models risks perpetuating the explanatory vacuum that current cranial osteopathy research inhabits.[12] Federated learning approaches may partially address the sample size problem while preserving participant privacy, but require harmonised data acquisition protocols that the field does not yet possess.[10]

A Research Roadmap:-

Translating the convergent potential of digital neurophenotyping and AI into scientifically credible cranial osteopathy research requires a phased, methodologically conservative approach. The following conceptual roadmap is proposed as a starting point for interdisciplinary discussion rather than a prescriptive protocol.

Phase I, Instrumentation Validation: Before outcome data from cranial osteopathic interventions can be meaningfully interpreted, the sensitivity of the proposed digital biomarker battery to known neurophysiological perturbations must be established.[6,9] This requires standardised validity studies demonstrating that selected EEG spectral indices, HRV metrics, and cognitive assessment tools respond predictably to interventions of known autonomic effect, such as paced breathing and progressive muscle relaxation, in healthy adults.[6,9]

Phase II, Proof-of-Concept Mechanistic Study: A within-person crossover design, with participants serving as their own controls across an active cranial osteopathic treatment session, a sham contact session matched for practitioner time, physical contact, and therapeutic attention, and a no-contact rest control, provides the minimum design necessary to detect treatment-specific physiological signals above non-specific effects.[2] Sample sizes of n=30-50 are feasible for this phase and sufficient to characterise effect size distributions for subsequent power calculations.[11]

Phase III, Adequately Powered Randomised Trial: Effect size estimates from Phase II inform a pre-registered, adequately powered randomised controlled trial incorporating blinded physiological outcome assessment. Primary endpoints should be pre-specified digital biomarkers from Phase I validation; patient-reported outcome measures serve as secondary endpoints.[7,10] Responder-subgroup analysis, guided by machine learning clustering of Phase II trajectories, enables stratified randomisation to maximise sensitivity without inflating type I error.[11] Priority research questions for this roadmap include: Does a single cranial osteopathic treatment session produce reproducible, measurable shifts in HRV frequency-domain indices relative to sham?[6] Are EEG alpha-band power or fronto-occipital coherence indices sensitive to cranial osteopathic intervention in a within-person repeated-measures design?[9] Do pre-treatment autonomic or sleep phenotypes predict physiological response magnitude?[6,8] Can machine learning clustering of multimodal neurophenotyping data identify subgroups for whom measurable change is consistently observed?[11] Affirmative or negative answers, rigorously obtained, would each constitute meaningful scientific progress.

Challenges and Cautions:-

Reproducibility. The replication crisis in biomedical research provides a sobering context for enthusiasm about novel measurement approaches.[10] Pre-registration of primary endpoints, open data policies, and multi-site validation studies are essential safeguards that must be embedded in the research culture of this field from its inception rather than retrofitted after initial positive findings.[7,10]

Signal interpretation. Even a robust, replicated shift in HRV or EEG power following cranial osteopathic intervention would not, by itself, validate the vitalistic mechanistic framework underpinning much practitioner discourse.[1] Signal changes require interpretation within established autonomic and cortical neuroscience frameworks, and mechanistic inference must be separated from efficacy claims.[6,9]

Data quality and participant burden. Continuous wearable monitoring imposes genuine participant burden, introduces artefact contamination from motion and electrode displacement, and raises compliance and dropout issues that are magnified in community-dwelling populations typical of osteopathic practice.[4,5]

Small effect sizes. If cranial osteopathic interventions produce neurophysiological changes, these are plausibly small in absolute magnitude and highly variable across individuals. Digital neurophenotyping may increase sensitivity, but cannot identify effects that are genuinely absent, and researchers must guard against post-hoc data dredging in high-dimensional datasets.[11,2]

Ethical considerations. The collection of continuous physiological and behavioural data raises substantive issues of informed consent, data security, secondary use governance, and algorithmic accountability.[10] Research protocols in this space must engage ethics review bodies with digital health expertise, particularly given India's Digital Personal Data Protection Act 2023.

Technological overreach. There is a risk that the availability of sophisticated measurement tools creates an unwarranted perception of scientific progress independent of the quality of the underlying research design. A poorly designed randomised trial with wearable sensors and a machine learning analysis module is not more rigorous than a well-designed observational study with validated questionnaires.[12] Methodological standards must not be lowered by enthusiasm for technology.

Conclusion:-

The scientific case for or against cranial osteopathic interventions cannot be adequately adjudicated with the methodological toolkit that has thus far been applied. This is not a trivial admission: it means that the existing literature, whether positive or negative in its conclusions, does not yet provide the evidentiary foundation necessary for confident clinical policy.[1,2] The emergence of digital neurophenotyping technologies and AI-assisted multimodal data analysis represents a genuine and potentially transformative opportunity to change this situation.[3,11,12] The opportunity is, however, contingent. It depends on the field committing to Phase I validation before clinical application, to pre-registered endpoints and open data policies, to mechanistic humility in the interpretation of positive findings, and to the active involvement of researchers from neuroscience, bioengineering,

and data science who bring both technical expertise and disciplinary scepticism.[7,10] The technology enables better questions; it does not guarantee better answers.

For the interdisciplinary research community, spanning AI, wearable sensing, neuroscience, and rehabilitation medicine, cranial osteopathy research represents a tractable and instructive test case for the application of digital neurophenotyping methodology to a domain characterised by contested mechanisms, poor measurement infrastructure, and genuine patient interest.[5,2] Regardless of what rigorous studies ultimately find, the process of building a scientifically credible measurement framework for manual medicine interventions will yield tools, methods, and insights of broad applicability across the emerging discipline of digital health.[10,12]

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