Short Communication

Debunking the myth of using “quiet” in clinical departments: an integrative overview of available literature

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Commonly referred to as the “art of treatment”, medical practice is an evidence-backed and peer-critiqued field of science. However, there exist several myths and misconceptions in current practice that seem to stem from personal idiosyncrasies and have been passed on over the years (telling and retelling) due to unavailability of logical explanation or credible evidence. Such idiosyncrasies provide the believers a sense of meaning, justification, and security [1].

In emergency medicine, there exists many such misconceptions including prevalence of “non-urgent cases”, and the influence of using words like “quiet” and “peaceful” on the levels of the crowding in the department [2,3]. In the present study, henceforth, we sought to investigate the latest evidence supporting the relationship between using the word “quiet” and its impact on clinical department overcrowding.

A PubMed database search from January 2015 to July 2022 for modern and current evidence-based English language articles (randomized controlled trials on human participants only) was conducted that investigated the relationship between the use of the word “quiet” and its corresponding impact on the perception of overcrowding in various clinical departments. Randomized Controlled Trials (RCTs) were included since they are considered the gold standard for comparing the effectiveness of various intervention strategies, given their ability to prevent bias in clinical trial design whilst controlling the efficacy of the statistical tests [4].

We were able to retrieve five such open-access full-text articles [2,3, 5–7], which we then benchmarked against the Joanna Briggs Institute’s (JBI) 2017 Critical Appraisal checklist for Randomized Controlled Trials (available at https://jbi.global/sites/default/files/2019-05/JBI_RCTs_Appraisal_tool2017_0.pdf; accessed 25th June 2022) for assessing the quality of the data reported (Table 1).

Most studies (four out of five) reported that usage of the term “quiet” was not associated with increased patient volumes or the perception of increased patient volumes. Although the findings may provide new evidence to debunk the myth and focus physicians’ attention on their work, a lack of rigorous methodology and lapses in adherence were identified in the included studies. The perceived strength of the available evidence is low for most studies (Table 1).

Systematic issues confounding all the investigated studies related to the participants’ repeated measures, which may impact the study’s conclusion, and the lack of a random allocation procedure because the participants may be in the intervention or control groups simultaneously. Furthermore, as the intervention is measured repeatedly on the same participants, blinding to the participants may fail, mainly due to the intervention of using the term “quiet” being a common myth for emergency practitioners [8]. As a result, blinding would not occur continuously throughout the study. To overcome this issue only Kuriyama et al. conducted sensitivity analysis and found no significant differences in the results of primary measured outcomes for both of their trials [5]. Another factor affecting the validity of the investigated studies is the non-matching of weekends and weekdays.

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Table 1
JBI’s Critical Appraisal checklist for RCTs.

| Criteria                                                                 | Kuriyama et al., 2016 [7] | Lamb et al., 2017 [5] | Brookfield et al., 2019 [4] | Go et al., 2022 [8] | Geller et al., 2022 [9] |
|-------------------------------------------------------------------------|----------------------------|-----------------------|-----------------------------|---------------------|------------------------|
| Was true randomization used for assignment of participants to treatment groups? | Yes                        | Yes                   | Yes                         | Yes                 | Yes                    |
| Was allocation to treatment groups concealed?                           | Yes                        | No                    | No                          | No                  | No                     |
| Were treatment groups similar at the baseline?                          | Yes                        | Yes                   | Yes                         | Yes                 | Yes                    |
| Were participants blind to treatment assignment?                        | Yes                        | Yes                   | No                          | Yes                 | Yes                    |
| Were those delivering treatment blind to treatment assignment?           | No                         | No                    | No                          | No                  | No                     |
| Were outcomes assessors blind to treatment assignment?                  | No                         | No                    | No                          | No                  | No                     |
| Were treatment groups treated identically other than the intervention of interest? | Yes                        | Yes                   | Yes                         | Yes                 | Yes                    |
| Was follow up complete and if not, were differences between groups interim of their follow up adequately described and analysed? | N/A                        | N/A                   | N/A                         | N/A                 | N/A                    |
| Were participants analysed in the groups to which they were randomized?  | N/A                        | N/A                   | N/A                         | N/A                 | N/A                    |
| Were outcomes measured in the same way for treatment groups?            | Yes                        | Yes                   | Yes                         | Yes                 | Yes                    |
| Were outcomes measured in a reliable way?                               | Yes                        | Yes                   | Yes                         | Yes                 | Yes                    |
| Was appropriate statistical analysis used?                               | Yes                        | No                    | Yes                         | Yes                 | No                     |
| Was the trial design appropriate, and any deviations from the standard RCT design (individual randomization, parallel groups) accounted for in the conduct and analysis of the trial | Yes                        | Yes                   | Yes                         | Yes                 | Yes                    |
| Additional Comments                                                     | Country of Trial           | Department of Investigation | Number of days investigated | Method of Randomization | Measurement of outcome of interest (workload reduction) |
|                                                                         | Japan                      | Emergency Department 347 shifts in two separate trials | Multi-centre 42 shifts | Computerized random number generator | Number of patient, stress levels, mealtime duration, fatigue, number of all admissions during their shift, and number of all ambulatory and transferred patients who visited the ED during their shift. |
|                                                                         | United Kingdom             | United Kingdom Clinical Microbiology 61 days | Big stick procedure | Number of referrals on a day | Number of visits and phone calls on a day |
|                                                                         | United States of America   | ENT Department 80 shifts | Microsoft® Excel 2016 Online randomizer | Number of consults per day and subjective survey on the perception of workload | Number of patient visits and workload (VAS scale) |
|                                                                         | United States of America   | Emergency Department 47 shifts Online randomizer | Matched groups, Ethics approval reported | Convenience sampling, multiple collection, exact participants not mentioned, subjective, no validation of VAS, single centre | Ethics approval reported |
|                                                                         |                             |                        |                            |                     |                        |
| Strengths of the study design                                           | The only registered trial  | Multi-centre study design | Inclusion of possibly affecting day: full moon, solstices or equinoxes, or a Friday on the 13th day of the month | Matched groups, Ethics approval reported |                        |
| Limitations of the study design                                         | Single centre, single blind, non-matched, no ethics reported | No ethics reported, non-matching amount of day, no weekday vs. weekend comparison, multi-centre but not evenly distributed, single blind | Non-matching amount of day, non-stratified randomization, no ethics reported, convenience sampling, single centre | Loss of data (8/80, 10%), depends on registrar note |                        |
| Use of appropriate reporting guidelines                                  | –                          | –                     | SPARR guidelines            | –                   |                        |
| Main Conclusion                                                          | No significant difference between control and intervention group | Significant differences between intervention and | No significant difference between control and intervention group | No significant difference between | No significant difference between |
|                                                                         |                            | intervention group     |                            | intervention group  |                        |
(continued on next page)
Baseline variations arising from personal beliefs of the investigated staff, their experience, knowledge on the topic, age, gender, and seniority have also largely been ignored in the reported studies. Lack of adherence to presenting proper reporting guidelines in three of the five studies also questions the reliability of these trials. Furthermore, when designing a randomized controlled trial, the convenience sampling method described in the studies is still debatable. Additionally, as the studied shifts/days are less than 100 in most of the studies, the different randomization methods employed by the studies might seem to be unsuitable as it can lead to unequal sample size between each group [9].

Finally, as previously indicated, four studies reported inconsequential findings, in contrast to one study that demonstrated significant results of using the phrase “quiet” in the multicentre orthopaedic emergency setting. However, the abundance of red flags in this study greatly raises doubts about how the findings might be interpreted. Firstly, utilizing a coin flip approach for allocation concealment is not ideal because it might be redone at the employee’s request and could result in an imbalance in baseline characteristics. As a result, it needs to be rethought before replication in a small-scale clinical trial [10]. Secondly, given the lack of valid excuse for the selection of shifts and the determination of sample size, the slightly significant p-value of 0.04 raises serious concerns about “p-hacking,” causing the underreporting of real effect sizes in published research.

In real-life scenarios, however, several tangible factors contribute to clinical department overcrowding - external (like patient demographics, accessibility, and devastating phenomenon) and internal factors (personnel availability, admission system, patient flow management, and hospital space). Overcrowding caused by these phenomena can have a significant impact on the economics of the hospital, psychology of the patients, and quality-of-care provided by the personnel (Fig. 1). Based on our review of the studied literature, there is no evidence that usage of any words or expectations of future clinical workload in emergency settings, including the use of “quiet” words, may influence departmental overcrowding. As a result, it was not included in our proposed cause-effect model.

Moving forward, future study designs should try to eliminate as many of the highlighted limitations of the previous studies, especially by conducting multicentre trials (which can offer larger sample sizes and larger observation periods for more generalizable findings). The published studies, despite their limitations, repetitively provide empirical support for the absence of an effect. Adequate reporting and publication of negative findings are informative because it allows researchers to deny a theory. Consequently, sufficient publication of non-significant findings makes scientific literature more complete, allowing for a more accurate assessment of a scientific work’s replicability.

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**Author contribution**

TPU and NJ conceptualized the report, whilst both authors were involved in the data collection and preparation of the manuscript. Both authors have read and agreed to the final version of the report for publication.

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None declared.

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