Correlation between disability measure and Black Hole (T1 hypointense lesion) lesion load in brain magnetic resonance imaging of patients with multiple sclerosis: Protocol for a systematic review and meta-analysis

Amir Valizadeh  
Tehran University of Medical Sciences  
https://orcid.org/0000-0001-5983-8527

Elham Barati  
Tehran University of Medical Sciences  
https://orcid.org/0000-0002-1750-2731

Mohammad Ali Sahraian  
Tehran University of Medical Sciences  
https://orcid.org/0000-0002-3224-8807

Mohammad Reza Fattahi  
Tehran University of Medical Sciences  
https://orcid.org/0000-0002-0770-0654

Mana Moassemi (Moassemi@gmail.com)  
Tehran University of Medical Sciences, Building no.1, Northern gate of the university  
https://orcid.org/0000-0002-0111-7791

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Abstract

Rationale: As the role of neurodegeneration in the pathophysiology of multiple sclerosis (MS) has become more prominent, the formation and evolution of chronic or persistent T1-hypointense lesions (Black Holes) have been used as markers of axonal loss and neuronal destruction to measure disease activity. However, findings regarding this subject are controversial. In this study we aim to clarify the level of importance of T1 hypointense lesions for estimating the prognosis of patients.

Objectives: To evaluate the correlation between T1 hypointensities (Black holes) lesion load (lesion mean volume) on brain MRI with disability level of patients with Relapsing-Remitting Multiple Sclerosis (RRMS) or Secondary-Progressive Multiple Sclerosis (SPMS).

Data sources: We will search MEDLINE (through PubMed), Embase, CENTRAL, Science Citation Index – Expanded (Web of Science), and Conference Proceedings Citation Index – Science (Web of Science). We won’t consider any timeframe, language, or geographical restrictions.

Methods: Standard systematic review protocol methodology is employed. Eligibility criteria is reported in line with PICOTS system. Population is limited to adult patients diagnosed with RRMS or SPMS, based on the McDonald criteria. Index (prognostic factor) of interest will be T1 hypointense (black hole) lesion mean volume (lesion load) on brain Magnetic Resonance Imaging (MRI). There will be no comparators. Outcome of interest will be the disability measure using Expanded Disability Status Scale (EDSS). For the timing domain, we will include studies only if the outcome was measured at the same time MRI was performed (or with a very close time interval between). Inpatient and outpatient settings will both be included. All included studies will be assessed for the risk of bias using a tailored version of the Quality In Prognosis Studies (QUIPS) tool. Extracted correlation coefficients will be converted to the Fisher's z scale and a meta-analysis will be performed on the results. We will then convert back the results to correlation coefficients again for the sake of presentation. For the purpose of assessing heterogeneity we will use prediction intervals. If feasible, we will also try to perform subgroup and sensitivity analyses. We will also evaluate the publication bias using Funnel plots and assess the confidence in cumulative evidence using an adapted version of the GRADE for prognostic factor research.

Introduction

Rationale

Multiple sclerosis (MS) is a chronic, disabling disease that targets patients in work productive years. Prognostic factors play important roles in evaluating the cost of the disease on both individual patients and the social economy. Since there is no exact method to predict MS progression confidently, four general MS phenotypes have been defined: clinically isolated syndrome (CIS), relapsing-remitting MS (RRMS), secondary progressive (SPMS), and primary progressive (PPMS) [2, 3]. RRMS is the most common MS phenotype. It is characterized by noticeable exacerbation or initiation of neurologic symptoms. These attacks, also called relapses are followed by periods of recovery, called remissions.
Some patients who are diagnosed with RRMS will gradually convert to a secondary progressive course which is a progressive deterioration of neurologic function over time. The other two types are not mentioned in this study.

Magnetic resonance imaging (MRI) is a sensitive paraclinical test for diagnosis and assessment of disease progression in MS and is often used to evaluate therapeutic efficacy. As the role of neurodegeneration in the pathophysiology of MS has become more prominent, the formation and evolution of chronic or persistent T1-hypointense lesions (black holes) have been used as markers of axonal loss and neuronal destruction to measure disease activity. On the one hand, T1 lesion load showed greater cross-sectional [4] and longitudinal [4, 5] correlations with Expanded Disability Status Scale (EDSS[6]) scores (a scale extensively used in studies for the assessment of disability for patients with MS) for patients with RRMS or SPMS than did T2 lesion load. On the other hand, several other studies reported an absence of correlations between T1 hypointensity and EDSS in patients with SPMS [7-11] and RRMS [9, 12].

In this study, we aim to review all the relevant studies, to evaluate the correlation between T1 hypointensities (Black holes) on brain MRI with disability level of the patients with MS. Our main goal is to clarify if this possible prognostic factor can be used to help to determine the prognosis of patients.

Objectives

To evaluate the correlation between T1 hypointensities (Black holes) lesion load (lesion mean volume) on brain MRI with disability level of patients with RRMS or SPMS.

Methods

Design and methods used for this protocol comply with Centre for Reviews and Dissemination (CRD’s) Guidance For Undertaking Reviews in Healthcare [13] and is reported in line with Preferred Reporting Items for Systematic Reviews and Meta-Analyses Protocols (PRISMA-P) [14]. Eligibility criteria were informed using the PICOTS [15] system.

Eligibility criteria

(P) Population: adult patients diagnosed with Relapsing-Remitting Multiple Sclerosis (RRMS) or Secondary Progressive Multiple Sclerosis (SPMS), based on the McDonald criteria [2, 16-18]:

- Two attacks or symptom flare-ups (lasting at least 24 hours with 30 days between attacks), plus two lesions.
- Two attacks, one lesion, and evidence of dissemination in space (or a different attack in a different part of the nervous system).
• One attack, two lesions, and evidence of dissemination in time (or finding a new lesion — in the same location — since the previous scan, or presence of immunoglobulin, called oligoclonal bands in the spinal fluid).

• One attack, one lesion, and evidence of dissemination in space and time.

• Worsening of symptoms or lesions and dissemination in space found in two of the following: MRI of the brain, MRI of the spine, and spinal fluid.

• **Relapsing-Remitting course**: A multiple sclerosis course characterized by relapses with stable neurological disability between episodes.

• **Progressive course**: A multiple sclerosis course characterized by steadily increasing objectively documented neurological disability independent of relapses. Fluctuations, periods of stability, and superimposed relapses might occur. Primary progressive multiple sclerosis (a progressive course from disease onset) and secondary progressive multiple sclerosis (a progressive course following an initial relapsing-remitting course) are distinguished.

(I) **Index (Prognostic factor)**: T1 hypointense (black hole) lesion mean volume (lesion load) on brain Magnetic Resonance Imaging (MRI)

(C) **Comparator**: not applicable

(O) **Outcome**: disability measure using Expanded Disability Status Scale (EDSS)

(T) **Timing**: measured at the same time MRI was performed (or with a very close time interval between)

(S) **Setting**: any

**Information sources**

The search will employ sensitive topic-based strategies designed for each database with no time frame limitations. There will be no language or geographical restrictions either. We will perform our search on the 10th of February, 2021.

**Databases:**

- MEDLINE through PubMed (RRID:SCR_004846)
- Embase (RRID:SCR_001650)
- Cochrane Central Register of Controlled Trials (CENTRAL) (RRID:SCR_006576)
- Science Citation Index –Expanded (Web of Science) (RRID:SCR_017657)
- Conference Proceedings Citation Index – Science (Web of Science) (RRID:SCR_017657)
Search strategy

Our search strategies for all the databases included in our study, namely MEDLINE (through PubMed), Embase, Science Citation Index – Expanded (Web of Science), and Conference Proceedings Citation Index – Science (Web of Science) are presented in appendix A.

Study records

Data management

Records will be managed through EndNote (RRID:SCR_014001) version X9 [19]; specific software for managing bibliographies.

Selection process

Two reviewers (AV and MM) will independently screen the title and abstract of identified studies for inclusion. We will link publications from the same study to avoid including data from the same study more than once. If any study cannot be clearly excluded based on its title and abstract, its full text will be reviewed. A study will be included when both reviewers independently assess it as satisfying the inclusion criteria from the full text. A third reviewer (MF) will act as arbitrator in the event of disagreement following discussion. We will prepare a flow diagram of the number of studies identified and excluded at each stage in accordance with the PRISMA flow diagram of study selection [1].

Data collection process

Using a standardized form, two reviewers (AV and MM) will extract the data independently. We will resolve any disagreements by discussion or, if required, by consultation with a third review author (MF). We will attempt to extract data presented only in graphs and figures whenever possible but will include such data only if two reviewers independently obtain the same result. If studies are multi-center, then where possible we will extract data relevant to each. In the case of missingness of data, if possible, we will try to contact the original investigators to request missing information. In case that was unsuccessful, we will only analyze the available information and will not try to impute any missing data.

Data items
Data extracted will include the following summary data: sample characteristics, sample size, study methods, inclusion and exclusion criteria, MRI settings used, founding sources, declarations of interests, and results.

Outcomes and prioritization

Our main outcome of interest is the relationship between participants’ EDSS score and T1 hypointense lesion mean volume.

Risk of bias in individual studies

Two review authors (AV and MM) will assess the risk of bias of each included study. We will resolve any disagreements by consensus, or by consultation with a third review author (MF). Because at the moment there is no standard tool for assessing the risk of bias in overall prognosis studies, we will use a tailored version of the Quality In Prognosis Studies (QUIPS) tool for assessing the risk of bias in studies [20], presented in Appendix B. Our tailored version of the tool consists of six risks of bias domains: study participation, study attrition, prognostic factor measurement, outcome measurement, study confounding, and statistical analysis and reporting. The study participation domain consists of five items: an adequate description of the source population or population of interest, adequate description of the baseline study sample, adequate description of the sampling frame and recruitment, adequate description of the period and place of recruitment, and adequate description of inclusion and exclusion criteria. The study attrition domain consists of four items: description of attempts to collect information on participants who dropped out, reasons for loss to follow-up provided, an adequate description of participants lost to follow-up, and no important differences between participants who completed the study and those who did not. The prognostic factor measurement domain consists of two items: provision of clear definition or description of the prognostic factor, and reporting of continuous variables or use of appropriate cut points. The outcome measurement domain consists of three items: provision of a clear definition of the outcome, use of an adequately valid and reliable method of outcome measurement, and use of same method and setting of outcome measurement in all study participants. The study confounding domain consists of the seven items: measurement of all important confounders, provision of clear definitions of the important confounders measured, adequately valid and reliable measurement of all important confounders, use of same method, and setting of confounding measurement in all study participants, appropriate imputation methods used for missing confounders (if applicable), important potential confounders accounted for in the study design, and important potential confounders accounted for in the analysis. The statistical analysis and reporting domain consists of two items: sufficient presentation of data to assess the adequacy of the analytic strategy, and an adequate statistical model for the design of the study.

Data synthesis
We will use R version 4 [21] as the software for our data synthesis. We expect correlation coefficients (r) to be our primary outcome measure. Most meta-analysts do not perform syntheses on the correlation coefficient itself because the variance depends strongly on the correlation. Rather, the correlation is converted to the Fisher’s z scale and all analyses are performed using the transformed values [22]. For the meta-analysis of correlation data, we first convert the correlation coefficients to Fisher’s z scale. Then we will calculate the variance and standard error of the Fisher’s z. We will perform a meta-analysis on those values based on the random-effects model. Finally, we will convert back Fisher’s z to correlation coefficient (r) for the sake of presentation.

**Assessment of heterogeneity**

We expect some heterogeneity between studies because of ethnicity and methodological diversity. We will report the range of the effects of the random-effects meta-analyses using prediction intervals. In a random-effects meta-analysis, the prediction interval reflects the whole distribution of effects across study populations, including the effect expected in a future study [23, 24].

**Subgroup analysis**

If at least 5 studies are available for each classification of MS in our study (RRMS and SPMS), we will perform a subgroup analysis for each of them.

**Sensitivity analysis**

We plan to perform sensitivity analyses to explore the influence of the following factors:

- Studies at high or unclear risk of bias
- Very long or large studies to establish the extent to which they dominate the results.

**Meta-bias**

To evaluate the risk of reporting bias across studies, a test for funnel plot asymmetry will be conducted. This test examines whether the relationship between estimated effect size and study size is greater than chance [25]. Funnel plots will be generated for visual inspection of potential publication bias. In the presence of publication bias, the plot will be symmetrical at the top, and data points will increasingly be missing from the middle to the bottom parts of the plot [26].
Confidence in cumulative evidence

The strength of the overall body of evidence will be assessed using an adapted version of the Grading of Recommendations, Assessment, Development and Evaluation (GRADE) framework for prognostic factor research [27], which takes into account seven criteria: study limitations, inconsistency, indirectness, imprecision, publication bias, moderate/large effect size, and dose effect. Two review authors (AV and MM) rate the certainty of the evidence for the outcome as 'high', 'moderate', 'low', or 'very low'. We resolve any discrepancies by consensus, or, if needed, by arbitration by a third review author (MF).

Appendices

Appendix A

Embase

#8 #7 AND #1
#7 #6 AND #5
#6 disability:ab,ti,kw
#5 #2 OR #3 OR #4
#4 'black hole*':ab,ti,kw
#3 (t1 NEAR/5 hypointens*):ab,ti,kw
#2 (t1 NEAR/5 lesion*):ab,ti,kw
#1 'multiple sclerosis'/exp

Medline through PubMed

Multiple Sclerosis[MeSH Terms] AND ((disability[tiab]) AND (“T1 hypointens*”[tiab] OR "T1 lesion*”[tiab] OR "black hole*”[tiab]))

CENTRAL

#1 MeSH descriptor: [Multiple Sclerosis, Relapsing-Remitting] explode all trees
#2 MeSH descriptor: [Multiple Sclerosis, Chronic Progressive] explode all trees
#3 #1 OR #2

#4 (t1 NEAR/5 lesion*):ab,ti,kw

#5 (t1 NEAR/5 hypointens*):ab,ti,kw

#6 'black hole*':ab,ti,kw

#7 #4 OR #5 OR #6

#8 disability:ab,ti,kw

#9 #8 AND #7

#10 #9 AND #3

Web of Science

#1 TS="Multiple Sclerosis"

#2 TS=(t1 NEAR/5 lesion*)

#3 TS=(t1 NEAR/5 hypointens*)

#4 TS="black hole*"

#5 #2 OR #3 OR #4

#6 TS="disability"

#7 #6 AND #5

#8 #7 AND #1

Appendix B

Our tailored version of the Quality In Prognosis Studies (QUIPS)
| Signaling question | Authors' judgment for ‘yes’ |
|--------------------|----------------------------|
| **Study participation: yes/no/partly/unsure/NA** | |
| a. Adequate participation in the study by eligible individuals | NA: usually participants with information on T1 hypointensity in brain MRI and information on disability levels or relapse rate are selected from a greater study cohort |
| b. Adequate description of the source population or population of interest | The source population for the cohort with T1 hypointensity in MRI is clearly described |
| c. Adequate description of the baseline study sample | Age and number of participants at baseline is clearly described |
| d. Adequate description of the sampling frame and recruitment | Way of establishing the source population, selection criteria, and key characteristics of the source population clearly described |
| e. Adequate description of the period and place of recruitment | The period and place of recruitment for both baseline and follow-up visits are clearly described |
| f. Adequate description of inclusion and exclusion criteria | Criteria used for diagnosis and classification and description of other inclusion and exclusion criteria are clearly described |
| **Study participation: risk of bias rating (high/low/unclear)** | High: most items are answered with ‘no’; Low: all items answered with ‘yes’; Unclear: most items are answered with ‘unsure’ Note: potentially a single item may introduce a high risk of bias, depending on study specifics |
| **Study attrition: yes/no/partly/unsure/NA** | |
| a. Adequate response rate for study participants | NA: usually participants with information on T1 hypointensity in brain MRI and information on disability levels or relapse rate are selected from a greater study cohort |
| b. Attempts to collect information on participants who dropped out described | Attempts to collect information on participants who dropped out are described (e.g., telephone contact, mail, registers) |
| c. Reasons for loss to follow-up provided | Reasons on participants who dropped out are available (e.g., deceased participants between baseline and follow-up, participants moving to another location) |
| d. Adequate description of participants lost to follow-up | Key characteristics of participants lost to follow-up are described (T1 hypointensity load) |
| e. No important differences between participants who completed the study and those who did not | Study authors described differences between participants completing the study and those who did not as not important or information provided to judge the differences |
| **Study attrition: risk of bias rating (high/low/unclear)** | High: most items are answered with ‘no’; Low: all items answered with ‘yes’; Unclear: most items are answered with ‘unsure’ Note: potentially a single item may introduce a high risk of bias, depending on study specifics |
### Prognostic factor measurement: yes/no/partly/unsure/NA

| Item                                                                 | Rating                                      |
|---------------------------------------------------------------------|---------------------------------------------|
| a. Clear definition or description provided                        | Study authors clearly indicate how they measure T1 hypointense lesions |
| b. Adequately valid and reliable method of measurement              | NA: brain MRI is the only available method for measuring T1 hypointense lesions |
| c. Continuous variables reported or appropriate cut-points used     | The exact size of the T1 hypointense lesion is reported and it is not categorized base on cut points |
| d. Same method and setting of measurement used in all study participants | NA: brain MRI is the only available method for measuring T1 hypointense lesions |
| e. Adequate proportion of the study sample had complete data        | NA: usually participants with information on T1 hypointensity in brain MRI and information on disability levels or relapse rate are selected from a greater study cohort |
| f. Appropriate methods of imputation were used for missing data     | NA: missing radiological size of T1 hypointense lesion can't be imputed |

#### Prognostic factor measurement: risk of bias rating (high/low/ unclear)

- **High:** most items are answered with 'no'; **Low:** all items answered with 'yes'; **Unclear:** most items are answered with 'unsure'
- Note: potentially a single item may introduce a high risk of bias, depending on study specifics

### Outcome measurement: yes/no/partly/unsure/NA

| Item                                                                 | Rating                                      |
|---------------------------------------------------------------------|---------------------------------------------|
| a. Clear definition of the outcome provided                        | Measuring disability level using EDSS should be clearly defined |
| b. Use of an adequately valid and reliable method of outcome measurement | EDSS should be measured by a trained physician |
| c. Use of same method and setting of outcome measurement in all study participants | Measurements of EDSS are the same for all participants |

#### Outcome measurement: risk of bias rating (high/low/unclear)

- **High:** most items are answered with 'no'; **Low:** all items answered with 'yes'; **Unclear:** most items are answered with 'unsure'
- Note: potentially a single item may introduce a high risk of bias, depending on study specifics

### Study confounding: yes/no/partly/unsure/NA

| Item                                                                 | Rating                                      |
|---------------------------------------------------------------------|---------------------------------------------|
| a. Measurement of all important confounders                        | Important confounders are: ethnicity, comedications, and comorbidities, socioeconomic status |
| b. Provision of clear definitions of the important confounders measured | Measurement of confounders has to be clearly described |
| c. Adequately valid and measurement of confounders is valid and reliable | Measurement of confounders is valid and reliable |
| Reliable measurement of all important confounders |
|--------------------------------------------------|
| d. Use of same method and setting of confounding measurement in all study participants |
| Measurements of confounders are the same for all study participants |
| e. Appropriate imputation methods used for missing confounders (if applicable) |
| Strategy to impute missing confounder data is described |
| f. Important potential confounders were accounted for in the study design |
| Methods section of the publication describes the strategy to account for confounders |
| g. Important potential confounders were accounted for in the analysis |
| Important confounders are accounted for in multivariable logistic regression and Cox proportional hazards models |
| **Study confounding measurement: risk of bias rating (high/ low/unclear)** |
| High: most items are answered with ‘no’; Low: all items answered with ‘yes’; Unclear: most items are answered with ‘unsure’ Note: potentially a single item may introduce a high risk of bias, depending on study specifics |
| **Statistical analysis and reporting: yes/no/partly/unsure/NA** |
| a. Sufficient presentation of data to assess the adequacy of the analytic strategy |
| Mean or median values, including confidence intervals or standard errors or standard deviations |
| b. Strategy for model building is appropriate and based on a conceptual framework or model |
| NA: we do not anticipate conceptual frameworks or explicit model building strategies for this type of research question (focusing on one prognostic factor only) |
| c. Statistical model is adequate for the study design |
| Mainly correlation coefficients |
| d. No selective reporting of results |
| NA: EDSS score and relapse rate are the only outcomes; if missing the study will be excluded |
| **Statistical analysis and reporting: risk of bias rating (high/ low/unclear)** |
| High: most items are answered with ‘no’; Low: all items answered with ‘yes’; Unclear: most items are answered with ‘unsure’ Note: potentially a single item may introduce a high risk of bias, depending on study specifics |

**No:** no or no relevant information to answer the signaling question

**Partly:** information is provided but not completely

**Unsure:** not enough information to answer signaling question with yes or no

**NA (not applicable):** signaling question not appropriate for this type of prognostic review

**Declarations**
Contributions

AV is the leading author for protocol development, analyses, and dissemination. AV is also the first reviewer. MM is the corresponding author. All authors will contribute to data interpretation and article drafts.

Amendments

Important protocol amendments post registration will be recorded and included in dissemination.

Support

No sources of support or funding were provided for this review.

Conflicts of interest

All authors declare there are no conflicts of interest regarding this study or its possible results.

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