Multigenomics Reveals the Causal Effect of Herpes Simplex Virus in Alzheimer’s Disease: A Two-Sample Mendelian Randomization Study

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In recent years, the herpes virus infectious hypothesis for Alzheimer’s disease (AD) has gained support from an increasing number of researchers. Herpes simplex virus (HSV) is a potential risk factor associated with AD. This study assessed whether HSV has a causal relationship with AD using a two-sample Mendelian randomization analysis model. Six single-nucleotide polymorphisms (SNPs) associated with HSV-1 and thirteen SNPs associated with HSV-2 were used as instrumental variables in the MR analysis. We estimated MR values of relevance between exposure and the risk of AD using inverse-variance weighted (IVW) method, MR-Egger regression (Egger), and weighted median estimator (WME). To make the conclusion more robust and reliable, sensitivity analyses and RadialMR were performed to evaluate the pleiotropy and heterogeneity. We found that anti-HSV-1 IgG measurements were not associated with risk of AD (OR, 0.96; 95% CI, 0.79–1.18; p = 0.736), and the same was true for HSV-2 (OR, 1.03; 95% CI, 0.94–1.12; p = 0.533). The findings indicated that any HSV infection does not appear to be a genetically valid target of intervention in AD.

Keywords: HSV, genome, Mendelian randomization, Alzheimer’s disease, causality

1 INTRODUCTION

Alzheimer’s disease (AD) is a complex chronic progressive degenerative disorder of the central nervous system, affecting primarily the elderly, which severely reduces the quality of life (Calabrò et al., 2021). According to the 2015 World Alzheimer Report, the number of AD patients is expected to double every 20 years, reaching up to 131.5 million by 2050 (Prince et al., 2015; Du et al., 2018) with the incidence rate of AD increasing exponentially after 65 years of age (Hou et al., 2019). AD is diagnosed after age 65 as late-onset AD (LOAD) and before age 65 as early-onset AD (EOAD). LOAD accounts for about 95% of AD cases. EOAD is essentially an inherited disease, with a 92%–100% heritability. In contrast, there are multiple factors influencing LOAD, which are sporadic (Laval and Enquist, 2021). AD has two central pathological features: the extracellular deposition of

Abbreviations: AD, Alzheimer’s disease; APOE-ε4, apolipoprotein e gene; CI, confidence interval; GWAS, genome-wide association studies; HSV, herpes simplex virus; IGP, International Genomics of Alzheimer’s Project; IV, instrumental variables; IVW, inverse-variance weighted; LD, linkage disequilibrium; MR, Mendelian randomization; OR, odds ratio; WME, weighted median estimator.
Amyloid plaques and intracellular accumulation of neurofibrillary tangles (NFTs). Amyloid plaques are mainly composed of amyloid-β (Aβ) protein and NFTs are composed of hyperphosphorylated tau proteins. Hence, there have been contrasting theories proposed about the underlying pathogenesis of AD, such as amyloid cascade hypothesis, Tau protein hypothesis, and oxidative stress. Nonetheless, to date, current therapies have failed to delay disease progression. In recent years, the herpes virus infection hypothesis has received a renewed interest by scientists who believe that infection is the main cause of AD.

In the 1980s, herpes simplex virus (HSV) was first proposed to be associated with AD after viral genetic material was discovered in the human brain as well as virus-induced lesions present in the limbic system were associated with AD (Ball, 1982). The viruses belong to the Alphaherpesviridae subfamily of the Herpesviridae family, including HSV-1 and HSV-2, which are ubiquitous human pathogens (Piret and Boivin, 2020). Previous studies (Wozniak et al., 2010) found that HSV-1 DNA was present in the brains of both AD patients and normal elderly people; however, in the brains of AD patients, HSV-1 DNA was found within 90% of the plaques and 72% of HSV-1 DNA was associated with plaques, while in the brains of normal elderly people, only 24% of HSV-1 DNA was associated with plaques. Thus, it was proposed that the HSV-1 infects infants and remains latent in the peripheral nervous system. Reactivation of latent HSV-1 infections may cause local neuronal damage and inflammation, which over time may lead to the deposition of Aβ and abnormal phosphorylation of tau in the brain. A recent study proposed that Aβ deposition and abnormal phosphorylation of tau were the brain’s immune response to HSV-1 (Eimer et al., 2018). However, another recent study showed that AD associated β-amyloid does not protect against HSV-1 infection in the mouse brain (Bocharova et al., 2021).

To date, the precise molecular events, and biological pathways underlying the disease have yet to be identified and the existing evidence does not definitively support the herpesviruses hypothesis of AD. The deposition of Aβ and abnormal phosphorylation of tau are not necessarily the cause of AD, but may be the result of other risk factors leading to AD. Meanwhile, given the existence of unmeasured confounding variables and reverse causation, previous epidemiological studies have demonstrated a correlation but no direct causal relationship between HSV and AD, which allows for a re-evaluation of the theory as a possible strategy.

Multi-omics research probes the interaction between multiple factors in biological systems, including genomics, epigenomics, transcriptomics, proteomics, metabolomics, and microbiomics. These factors jointly affect phenotypes and physiological traits. With the development of high-throughput sequencing technology, omics research continues to provide more extensive data. Through high-throughput sequencing, omics, and data integration studies, we can comprehensively and systematically understand the relationship between various factors in the fields of basic research, molecular biology, clinical diagnosis, and drug discovery. (Hasin et al., 2017).

Genomics is the earliest discipline stemming from histology, and focuses on the study of the entire genome, and is currently the most established discipline in the field. Genomics focuses on the identification of genetic variants associated with disease, treatment response, or patient prognosis (Hasin et al., 2017). With the successful development of next-generation sequencing (NGS) technology and the completion of the human genome project and the International Human Genome HapMap project (HapMap), genome-wide association studies (GWAS) have become a method for identifying millions of genetic variants related to complex diseases (GWAS catalog https://www.ebi.ac.uk/gwas/home) in different human populations. In such studies, millions of individuals are genotyped for many genetic markers, and the genotypes and phenotypes are subjected to statistical analysis at a population level. Significant differences in minor allele frequencies (MAF) between cases and controls are thought to be markers affecting the trait. GWAS studies provide an invaluable contribution to our understanding of complex phenotypes (Hasin et al., 2017).

Mendelian randomization (MR) is a strategy for evaluating the causality of risk factors of a disease using genetic variants from the GWAS as instrumental variables (IV) (Lawlor et al., 2008). It is based on the Mendelian inheritance law of “random allocation of parental alleles to offspring” in meiosis, which is equivalent to a randomized controlled trial using genotypes. MR analysis can remove the limitations of traditional epidemiology. As alleles were randomly allocated at conception, confounders cannot influence the result of the allocated alleles. Because the disease cannot alter genetic variants, reverse causation may be avoided. IVs should satisfy three major hypotheses (Figure 1), which have been widely described in recent studies (Liu et al., 2018; Liu et al., 2021).

1) The IV is associated with the exposure (γ ≠ 0, strong IVs).
2) The IV is not associated with the confounders (ϕ = 0).
3) The IV does not influence the outcome through some pathways other than the exposure (α = 0, no directional pleiotropy).

Scepanovic et al. (2018) measured quantitative IgG responses to HSV-1 and HSV-2 infection in humoral immunity to explore the influence of genetic factors on the variability of humoral responses. After genome-wide genotyping of single-nucleotide polymorphisms (SNPs) and imputation, they examined associations between genetic variants and HSV-1 and HSV-2 IgG and performed two genome-wide association analyses. The International Genomics of Alzheimer’s Project Consortium (IGAP) (Lambert et al., 2013) conducted a meta-analysis using genotyped and imputed data on four previously published GWAS datasets and obtained a novel genome-wide association analysis demonstrating the relationship of genetic variants with AD. In the present study, we used many SNPs of multi-genome association analysis as IVs to perform two-sample MR analysis (Gibran et al., 2018).

2 MATERIALS AND METHODS

2.1 Data Sources
The exposure risk factors considered in this study were HSV-1 and HSV-2. The genetic variations for both exposures were anti-
HSV-1 IgG measurement and anti-HSV-2 IgG measurement, which were downloaded from a GWAS study of Scepanovic et al. (2018), which was the basis of the summary data published in the NHGRI-EBI GWAS (https://www.ebi.ac.uk/gwas). The sample was derived from The French Milieu Interieur cohort, which was stratified by sex (500 men, 500 women) and age (200 individuals from each decade of life, between 20 and 70 years of age). The HSV-2 datasets contained 208 cases and 792 controls, and HSV-1 datasets contained 645 cases and 355 controls.

The summary data of AD derived from the International Genomics of Alzheimer’s Project Consortium (IGAP), which was a sizeable two-stage research based on GWASs of AD in 74,046 diseased and normal individuals of European ancestry (Lambert et al., 2013). In stage 1, the IGAP performed a meta-analysis of four previously published GWAS datasets containing 17,008 AD patients and 37,154 controls, using genotyped and imputed data on 7,055,881 SNPs. The outcome data from IGAP stage 1 results were from the study of Kunkle et al. (2019). Table 1 shows the detailed descriptions of IGAP stage 1 data.

### Table 1 | Description of consortium datasets for IGAP stage 1.

| Consortium | N (cases/controls) | Percent women (cases/controls) | Case mean AAO (s.d.) | Control mean AAE (s.d.) |
|------------|--------------------|-------------------------------|----------------------|------------------------|
| ADGC       | 10,273/10,892      | 59.4/58.6                     | 74.7 (7.7)           | 76.3 (8.1)             |
| CHARGE     | 1,315/12,968       | 63.6/57.8                     | 82.7 (6.8)           | 72.8 (8.6)             |
| EADI       | 2,243/6,017        | 64.9/60.7                     | 68.5 (8.9)           | 74.0 (5.4)             |
| GERAD      | 3,177/7,277        | 64.0/51.8                     | 73.0 (8.5)           | 51.0 (11.8)            |
| N          | 17,008/37,154      |                               |                      |                        |

AAO, age at onset; AAE, age at examination.

2.2 Methods

All the analyses were performed using R version 4.1.0 software.

2.2.1 Selection of Instrumental Variables

The most critical step in MR design is to identify suitable genetic variants as IVs. First, we extracted SNPs that had significant ($p < 1 \times 10^{-3}$) associations with HSV-1 and HSV-2. Then, we performed a linkage disequilibrium (LD) analysis to exclude mutual linkage SNPs and to discard non-biallelic SNPs. LD ($r^2 < 0.001$, $kb > 10,000$) was applied to select IVs of HSV-1 and HSV-2. The samples used to estimate the LD effect derived from individuals of European ancestry from the 1,000 Genome Project. Correlated SNPs in LD were excluded using the “clump_data” function of the “TwoSampleMR” R package. As a result, 7 SNPs were identified for HSV-1 and 13 SNPs for HSV-2.

2.2.2 Harmonize

A summary set can generate errors if the effect alleles for the SNP effects in the exposure and outcome datasets are different. We aligned the effect alleles for exposure and outcome based on reported effect alleles and effect allele frequencies using the “harmonise_data” function of the “TwoSampleMR” R package (Gibran et al., 2018). Furthermore, we used F-statistics (Bowden et al., 2016) to measure the strength of the selected IVs. If the F-statistic was more than ten, genetic variants were generally deemed to be a strong IV.

2.2.3 Mendelian Randomization

We conducted the MR analysis using inverse-variance weighted (IVW) regression analysis, MR-Egger regression analysis, and weighted median estimator (WME). IVW can provide accurate estimates when the IV satisfies the MR assumptions that there are no invalid IVs (Burgess et al., 2013). The mean effect estimate of IVW is derived from a random effect IVW meta-analysis of the Wald ratios (SNP-outcome associations divided by SNP-exposure associations).
TABLE 2 | SNPs significantly (p-value < 1 × 10^-5) and independently (I^2 < 0.001) associated with herpes simplex virus type 1 (HSV-1) infection (SNPs = 6).

| Exposure | SNP        | Effect allele | Other allele | EAF | Beta    | SE    | p-value | F-statistic |
|----------|------------|---------------|--------------|-----|---------|-------|---------|-------------|
| HSV-1    | rs10977313 | T             | G            | 0.11 | −0.12   | 0.02  | 2.97 × 10^-7 | 26.9        |
|          | rs1446553  | A             | G            | 0.25 | 0.07    | 0.02  | 9.85 × 10^-6 | 19.9        |
|          | rs34018815 | A             | G            | 0.10 | −0.11   | 0.02  | 8.35 × 10^-6 | 20.2        |
|          | rs58599785 | T             | C            | 0.16 | 0.08    | 0.02  | 4.91 × 10^-6 | 21.3        |
|          | rs78421079 | T             | C            | 0.08 | −0.12   | 0.03  | 6.49 × 10^-6 | 20.7        |
|          | rs8020017  | G             | A            | 0.40 | −0.06   | 0.01  | 9.85 × 10^-6 | 19.9        |

TABLE 3 | Association of six SNPs for HSV-1 infection with AD using MR with different methods.

| Outcome | SNPs | OR   | 95% CI  | p-value | Pleiotropy Intercept p-value | Heterogeneity Cochran’s Q-statistic p-value | I^2 |
|---------|------|------|---------|---------|-----------------------------|---------------------------------------------|-----|
| AD      | 6    | IVW  | 0.96    | [0.79–1.18] | 0.736 | 0.833 | 5.836 | 0.323 | 958 |
|         |      | WME  | 0.97    | [0.77–1.23] | 0.663 | 0.664 | 5.896 | 0.232 |     |
|         |      | MR-Egger | 0.79 | [0.32–1.99] | 0.663 | 0.664 | 5.896 | 0.232 |     |

3 RESULTS AND DISCUSSION

3.1 The Causality of HSV-1 and AD

After removing the palindrome SNP (rs1738233), six SNPs for HSV-1 infection were identified, which were significant (p < 1 × 10^-5) and independent (r^2 < 0.001). The F-statistics for the six SNPs were all more than 10, which indicated that all six IVs were strong instruments (Table 2).

Table 3: Figure 2 showed the estimated associations of HSV-1 risk factor with AD from MR analysis. Genetically predicted HSV-1 infection was not associated with AD risk using IVW (OR = 0.96, p = 0.736), WME (OR = 0.97, p = 0.833), and MR-Egger (OR = 0.79, p = 0.653). The MR-Egger intercept indicated no directional pleiotropy (intercept = 0.018, p = 0.694), suggesting that horizontal pleiotropy was unlikely to influence the IVW estimate. The F statistic was 0.958, indicating that relative bias did not materially affect the standard MR-Egger analysis. Cochran’s Q test showed no existence of heterogeneity of SNPs (Cochran’s Q-statistic = 5.83, p = 0.322), while RadialMR showed that there were no outliers in the six SNPs.

3.2 The Causality of HSV-2 and AD

Thirteen SNPs for HSV-2 infection were identified, which were both significant (p < 1 × 10^-5) and independent (r^2 < 0.001) (Table 4). The F-statistics for the thirteen SNPs were more than 10, which indicated that they were strong IVs.

Table 5: Figure 3 showed the estimated associations of HSV-2 risk factors with AD from the MR analysis. Genetically predicted HSV-2 infection was not associated with the AD using IVW (OR...
Meanwhile, no outliers were detected using RadialMR. GWAS summary statistics data from the French Milieu Intérieur heterogeneity of SNPs (Cochran

Furthermore, the Cochran’s Q-statistic indicated the existence of heterogeneity of SNPs (Cochran’s Q-statistic = 18.8, p = 0.04). Meanwhile, no outliers were detected using RadialMR.

4 DISCUSSION

We found that both HSV-1 and HSV-2 were not causally associated with an increased risk of AD using genetic variation as instrumental variables. Kwok and Schooling (2021) used the GWAS summary statistics data from the French Milieu Intérieur cohort, the United Kingdom biobank, and the US 23 and Me Study, pointing out that HSV-1 and HSV-2 were not associated with AD. SY et al. (2021) used the GWAS summary statistics data from the 23 and Me cohort, indicating the same result.

4.1 The Result of HSV and AD

Although the causality of the association is unclear, many studies have proven that HSV is not unrelated to AD. HSV-1 virus was detected in the brains of both AD patients and elderly normal people. However, most of the AD patients were APOE-ε4 gene carriers. The herpesvirus hypothesis proposes that HSV-1 enters the brains of APOE-ε4 carriers, where it remains a latent life with limited transcription and low protein synthesis. In response to immunosuppression, peripheral infection, and inflammation, HSV-1 reactivates, creating a combination of viral action and inflammatory effects that are poorly repaired by APOE-ε4 carriers, ultimately leading to the development of AD (Itzhaki, 2018). In addition, a recent study pointed out novel molecular mechanisms through which recurrent HSV-1 infection may affect neuronal aging, likely contributing to neurodegeneration (Napoletani et al., 2021).

We inferred that our results may have occurred mainly due to several reasons. The major reason is that reactivation after latent HSV-1 infection may be responsible for a pathogenetic mechanism of AD, and IgM is a marker of activation of primary infection. Our study used anti-HSV-1 IgG antibodies rather than IgM as a proxy for HSV-1 infection, implicitly demonstrating that previous HSV-1 infection is not associated with AD risk. Another reason is the speculation that HSV-1 infection is not a risk factor for cognitive decline but rather a phenomenon that co-occurs with neuro-inflammation or as a result of neuro-inflammation.

Meanwhile, we found that HSV-2 was not causally associated with an increased risk of AD using genetic variation as an instrumental variable. This is probably because that according to the available epidemiological observations, HSV-2 mainly invades the genitalia and the area from the waist down and is not associated with the brain.

Future studies should perform MR analyses using anti-HSV-1 IgM antibodies as an IV for HSV-1 infection. What we can conclude, however, is that AD is not simply a single factor disease caused by HSV, but that it encompasses complex disease mechanisms.
4.2 Advantages and Challenges of MR Analysis

In the investigation of risk factors for AD, traditional research methods present many challenges in discovering the cause of the disease. Observational studies can only demonstrate a correlation rather than causality between exposure and outcome due to confounding factors and reverse causality. Cohort studies can make causal arguments but waste time. Random control trials (RCT) are considered the gold standard for clinical diagnosis and have a solid causal view. However, when applied by researchers, they are difficult to practice due to medical ethics and the many limitations of the design process. For these reasons, MR analysis has become a more convenient and effective way of exploring the causal links between risk factors and AD.

The application of MR analysis in this study has several advantages. First, reverse causality can be avoided, and second, it can prevent the interference of confounding factors. MR analysis can also address situations where an intervention experiment cannot be performed because of ethical restrictions (Zheng et al., 2017). Our exposure data were obtained from a publicly available GWAS database published with credibility. Our outcome data derived from a study conducted by the IGAP with a large sample population.

Nonetheless, our study also has some limitations. First, our data samples were based on individuals of European ancestry, so the results are not representative of all races. Second, the sample size of the exposure data was not sufficiently large, leading to the low power of statistics and false negatives. However, a significant number of IVs can lead to high power but inevitable heterogeneity and pleiotropy of IVs. This is where the general challenge of MR.

5 CONCLUSION

We implemented a two-sample MR to demonstrate the causal relationship between HSV infection and AD risk. The SNPs were independent and strong instrumental variables, and the result was robust and reliable. Our findings indicated the negative association between any HSV IgG and AD. Further research is needed to investigate whether HSV IgM is correlated with AD, and whether HSV infections that co-occur with neuro-inflammation are more relevant.

DATA AVAILABILITY STATEMENT

Publicly available datasets were analyzed in this study. The AD data can be found here: https://www.niagads.org/datasets/ng00075. The HSV data can be found here: https://www.ebi.ac.uk/gwas/efotraits/EFO_0009349 and https://www.ebi.ac.uk/gwas/efotraits/EFO_0009350.

AUTHOR CONTRIBUTIONS

XZ contributed to the conception and designed the study. LL and ZX organized the database. YZ performed the statistical analysis and wrote the article. JQ supervised the project and acquired the funding. XZ reviewed the article. All the authors contributed to the article and approved the submitted version.

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