Original Article

The Effect of Statins in Epilepsy: A Systematic Review

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Background and Objectives: Statins are inhibitors of 3-hydroxy-3-methylglutaryl coenzyme A (HMG-CoA) reductase, used for the management of hypercholesterolemia and related atherosclerotic diseases. Several studies have indicated the neuroprotective effects of stating on several neuropathological conditions. However, the role of these medications in epilepsy is still unclear. The purpose is to evaluate and summarize the level of evidence on the efficacy of statins in neuronal hyperexcitability and the neuroinflammatory processes of epilepsy. Methods: A systematic review was performed. Eligibility Criteria: This review involved studies conducted in humans and nonhuman experimental models, covering the use of an inhibitor of HMG-CoA reductase, alone or accompanied by another medication, in epilepsy. Information Sources: A systematic literature search was performed in PubMed, Embase, Ebsco Host, Scopus, Science Direct, Medline, and LILACS. Risk of Bias: It was evaluated with the Newcastle-Ottawa Scale and the experimental studies were evaluated using the GRADE tool. Results: Twenty articles of the 183 evaluated were included. Sixteen studies were conducted in animal models and four studies in humans. Most studies in mice reported a reduction in epileptiform activity and reduction in systemic inflammation with the treatment of statins, potentially influencing epilepsy control. Few studies in humans were performed in the geriatric population with variable results (neuroinflammation, seizure prevention, cell death, prevention of kindling, increase in convulsive threshold, increase in latency, decrease in frequency of crisis, and reduction in mortality) related to reduction in the rate of hospitalizations, mortality, and prevention of epilepsy. Studies in mice found a decrease in interleukin-1 β (IL-1 β), IL-6, and tumor necrosis factor alpha and an increase in IL-10 and endothelial nitric oxide synthase. Conclusions: The possible antiepileptic mechanism of statins may be related to the reduction in neuroinflammation mediated by a decrease in pro-inflammatory cytokines and action in the nitrergic system. Further studies evaluating the impact of statins on seizure control are necessary.

Keywords: Brain, epilepsy, seizure, statins

INTRODUCTION

 \mathcal{E} pilepsy is a neurological disease generated by an abnormal brain electrical activity in certain regions of the brain that predisposes to recurrent unprovoked attacks. Worldwide, a prevalence of 1%–2% is estimated, affecting between 50 and 65 million people, with 50,000– 100,000 new cases per year. In addition, it is estimated that 30%–40% are refractory to seizure treatment.^[1-3] Epilepsy is considered a public health problem worldwide and is one of the most frequent neurological disorders.^[3]

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Statins are inhibitors of 3-hydroxy-3-methyl glutaryl coenzyme A (HMG-CoA) reductase, used for the management of hypercholesterolemia and related atherosclerotic diseases, such as coronary artery

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disease.^[1,4,5] The enzyme HMG-CoA reductase catalyzes the conversion of HMG-CoA to L-mevalonate; statins prevent the biological activities of L-mevalonate due to the aforementioned inhibitory effect.^[5] Statins perform pleiotropic actions on the endothelium, the inflammatory response, or the production of free radicals. The inhibition of obtaining endogenous cholesterol induces a positive regulation of low-density lipoprotein (LDL) receptors on the cell surface. Obtaining a higher absorption of LDL from the blood and therefore a decrease in its concentration. In addition, high-density lipoprotein levels increase and triglyceride levels decrease.^[6]

Several studies have indicated the neuroprotective effects of statins on several neuropathological conditions. The anticonvulsive activity has been explained by suppressing reactive astrogliosis with neuroinflammation in the crises; by stimulating GABAergic activity and inhibiting glutamatergic; by modulating the glycogen synthase kinase- 3β pathway; and by decreasing the infiltration of monocytes in the neuronal death of the hippocampus and pro-inflammatory gene expression.^[1,4]

The lipophilic statins (atorvastatin [ATV], lovastatin, fluvastatin, pitavastatin, and simvastatin) can passively pass through the blood-brain barrier (BBB); in addition, the hydrophilic statins can also enter the neuroparenchyma.^[6,7] All statins are substrates for organic anion transporter polypeptides (OATPs), of which OATP1A2 and OATP1C1 are expressed in the brain. Despite this, the selectivity of hydrophilic statins for these subtypes has not been explored to determine their mechanism of entry to the central nervous system (CNS). In addition, the existence of monocarboxylic acid transporters in the BBB can constitute an alternate route of entry to the CNS although there are no specific studies for the CNS either. Independently of the specific transporters, it is feasible that statins are deposited at different speeds and concentrations within the CNS according to their different lipid solubility alone.[7]

Methods

Objectives

The objective of this review is to answer the following question: What is the level of evidence on the efficacy of statins to decrease neuronal hyperexcitability and neuroinflammatory processes of epilepsy?

To develop the review, the steps of the patients, intervention, comparison, outcomes, and study design strategy were followed.

Inclusion criteria

Types of participants

This review involved studies conducted in humans and nonhuman experimental models.

Type of intervention

It covered the use of inhibitors of HMG-CoA reductase, alone or accompanied by another medication.

Types of studies

The qualitative component of the review included studies that described the molecular mechanisms of statins in the CNS, the pharmacokinetic aspects, and the changes derived from the use of these drugs in neurotransmitters and cerebral cholesterol. The quantitative component included randomized and nonrandomized controlled trials, quasi-experimental designs, prospective or retrospective cohort studies, and nested case–control studies.

Types of results

The benefit of the therapy was defined as the decrease in neuroinflammation derived from seizures, decrease in neuronal death after the seizure, prevention of seizures, decrease in mortality after a seizure episode, increase in seizure threshold, reduction in the frequency, and increase in the latency of the crises.

Search and selection of studies strategy

This systematic review followed the recommendations of the Cochrane Collaboration (PRISMA). A bibliographic search was carried out in the databases: PubMed, Embase, Ebsco Host, Scopus, ScienceDirect, Medline, and LILACS, considering all the publications made up to February 2, 2018. The search was carried out in five steps: First, the keywords using the Medical Subject Headings and DeCs (Health Descriptors), then proceeded to use the descriptors of the subject (hydroxymethyl glutaryl-CoA reductase inhibitors, statins, epilepsy, epileptogenesis) in the databases mentioned. Subsequently, the duplicate records were eliminated (Step 2) and an analysis of the titles and abstracts thrown by the search was continued (Step 3), then the full-text review of the selected articles was proceeded (Step 4), and finally, the references were reviewed of the included articles to identify those studies that also met the eligibility criteria (Step 5). Letters to the editor and studies with a language other than English or Spanish were excluded.

Method of revision

The search strategy and selected studies were evaluated by two independent reviewers (LM and ZC). The discrepancies were discussed with a third reviewer (RM).

Data collection

The following data were extracted from the studies that met the eligibility criteria: authors, year of publication, number of participants, type of study, study objective, statin used, statin dose, epilepsy inducer used, route of administration of the statin, duration of treatment, and results of the study. These data were compiled in Microsoft Excel and were divided into two: collection of human studies and collection of non-human studies.

Assessment of quality and risk of study sessions

The assessment of the risk of bias in human studies was performed using the Newcastle–Ottawa Scale [Figures 1 and 2], while experimental studies in animal models were evaluated using the GRADE tool [Figure 3].

Data synthesis and additional analysis

An analysis of the studies was carried out in nonhuman experimental models that only involved the use of statins, excluding from this analysis those studies that applied a drug or substantial addition to the statin and the studies that did not specify the duration of the treatment. With the Epi-info 7.2 programs, (CDC, Atlanta, Georgia, USA) the median duration of treatment, absolute and relative frequency of the variables were determined: statin used, statin dose, epilepsy inducer used, and route of administration.

RESULTS

Selection of studies

The selection process of the studies was based on the PRISMA foundations as shown in Figure 4. Twenty articles of the 183 included in the bibliographic search were considered.

Characteristics of the studies

Studies in animal models

We identified 16 studies in animal models (mice), of this total only 10 studies (n = 626 mice) met the criteria for descriptive analysis [Table 1]. We excluded three studies that used an additional substance to the statin; one study that did not clearly specify the duration of the treatment and two studies that did not clarify the number of participants. The median duration of treatment with statins was 14 days (IQR = 7–15), the inducers of epilepsy used were pilocarpine, kainic acid, quinolinic acid, pentylenetetrazole (PTZ), and electroshock test, as shown in Figure 5. Sehar *et al.*'s study was performed in two phases: acute and chronic, for statistical analysis were taken as two studies. On the other hand, as can be seen Figure 6, the most used route of administration was



Figure 1: PRISMA flow diagram of our search mechanism

oral. In addition, the most commonly used statin dose was 10 mg of ATV [Figure 7]. In Table 2, it is observed that most of the studies reported a positive change of statin treatment.

Human studies

We identified four studies in humans (n = 1,071,422): 3 (75%) of retrospective cohort and 1 (25%) of nested cases and controls. The results are shown in Table 3.

Evaluation of the risk of bias

The results of the evaluation of the bias are shown in Figures 2-4.

Statins and brain cholesterol

One study quantified sterol levels in the hippocampus of an animal model with unilateral hippocampal lesion induced by kainic acid. The lovastatin treatment was performed 3 days before and 3 days after the induction of status epilepticus (SE). There were no significant changes in the levels of sterols (lanosterol and desmosterol) at 24 h after the SE, but it did detect a bilateral reduction in the hippocampus of the metabolite 24-OHC and cholesterol levels at 48 h and 14 days (P < 0.001 and P < 0.01, respectively). These periods in which changes were identified corresponded to the preepileptogenic and to the appearance of daily seizures. On the other hand, lovastatin was not associated with alteration of seizures during status.^[8]







Figure 4: Risk of bias using Newcastle–Ottawa Scale for case–control studies

Statins and neuroinflammation

We identified three studies related to the role of statins in neuroinflammation mediated by cytokines. Two studies used lovastatin and one study used ATV. All three studies administered postinduction statin with pilocarpine. The study by Oliveira et al. found a reduction of the pro-inflammatory cytokines: interleukin-1 β (IL-1 β), IL-6, tumor necrosis factor alpha (TNF- α), and interferon- γ in cortex and hippocampus, 14 days after induction (P < 0.05) together with an increase in the levels of the anti-inflammatory cytokine IL-10. These results were better with the dose of ATV 100 mg/kg compared to the dose of 10 mg/kg. In addition, the control group of ATV without induction with pilocarpine also showed a reduction in pro-inflammatory cytokines. In this same study, it was found that mice needed higher doses of the statin to achieve an increase in IL-10, unlike rats, which in turn had higher levels of pro-inflammatory cytokines.^[3] Gouveia et al.found similar results in both 2011 and 2014.^[5,9] The two



■ Low risk ■ High risk **Figure 3:** Risk of bias using Newcastle–Ottawa Scale for cohort studies





Percentage of studies



					Table 1: Stati	n studies in anim	als		
Authors	Year	Number	Statin	Number of	Inductor	Start of statin	Duration	Evaluations	Results
		of		times per			of		
		animals		day		-	reatment		
Sehar	2015	24	Controls=6	Once a day	PTZ (60 mg/kg)	1 h before PTZ	7 days	ICES test, elevated plus	ATV 80 mg/kg increased seizure
<i>et al</i> . ^[1]			ATV 20 mg, 40 mg			administration		maze and forced swim test	threshold current significantly
			and 80 mg=6 for						compared with the control group
			group						(22±0.00); aid not significantly increase the letenant of minutes
		60	Control 6-10		DT7 (75 ma/ha)	1 h hafara DT7	7 moolee	للمتعامدة ومنام	ATV 20 40 and 80 matrice
		00			L 1 Z (Z) 111 Z VZ)	1 II UCIUIC F 12	/ WCCKS	Nacille Scale allu	AIV 20, 40, and ov mg/ng
			Only F1Z=12 ATV 20 mg + PTZ			authinishauon		assessment of uppairme, glutamate, and GABA	of PTZ kindling significantly
			ALV 20 mg I 12, A0 mg + DT7 v					levels	in a dose-dependent manner
			80 mg + PTZ=12						(P<0.05, P<0.01, and P<0.001,
			for group						respectively)
Van Vliet	2011	13+	Control with	Once a day	ES	7 before the	14 days	Video-EEG	ATV did not affect the duration
<i>et al</i> . ^[18]			ATV=5, ATV 10			induction of		monitoring-Racine	of SE or the development
			mg/kg + ES=8,			epilepsy and 7		scale, blood-brain	of epilepsy, had not reduced
			vehicle+ ES=8			days after		barrier permeability,	inflammation, neuronal death, or
								and fluorescent	synaptic reorganization
		20		-			-		
Hevenn	7117	96	4 mg/kg	Unce a day	KA (0.5 µg)	1 n berore and 1	/ days	EEU telemetry units,	LV I did not alter seizures during SE
et al.						n atter the KA		sterol analysis, and	or seizure-induced neuronal death.
								histopathology	Changes in hippocampal cholesterol
									homeostasis occur bi-laterally
Uliveira	2018	108	Saline=22, AI V	Once a day	Pilocarpine	3 h atter	14 days	Cytokine analysis and	AIV dose-dependently decreased
$et al.^{[s]}$			10=17, ATV		(100 mg/kg) after	diazepam		Behavioral tests	basal and SE-induced levels of
			100=13		methylscopolamine	injection			$(\text{IL-1}\beta), (\text{IL-6}), (\text{TNF-}\alpha), (\text{INF-}\gamma)$
					1 mg/kg				and increased (IL-10) levels in the
									hippocampus and cerebral cortex
Gouveia	2014	26	Control=5; LVT	Twice daily	Pilocarpine	After 2 h of SE	15 days	Real-time PCR analysis	LVT induced an increased
lel al.			(20 mg/kg)=5;		(350 mg/kg)	onset		tor cytokines and	expression of the IL-10 and the
			Pilocarpine=8 and					Immunohistochemistry	pilocarpine plus lovastatin group
			rucarpine pius						showed a significant decrease in the leviale of II -18 and TME ~
			LV 1-0						during the letent and chronic
									duting arc faicht and chronic phase
Diermartiri	000	71	Saline + $OA=77$	Once a dav	Oninolinic Acid	Refore OA	7 davs	I -[3H] olutamate []'ntake	ATV 10 mo/kg nrevented seizures
$et al.^{[10]}$	1007	1/	ATV 1 mg/kg +	Olive a day	(36.8 nmol)	administration	o day o	Cell death (Propidium	induced by QA in 29.41% of the
			QA=15		~			Iodide Staining)	mice. $(P=0.004)$, ATV prevented
			ATV 10 mg/kg +						QA induced cell death in the
			QA=34						hippocampus and prevented the
									reduction in glutamate uptake into the hippocampus
									and seens a difference of the see
									Contd

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					Tabl	le 1: Contd			
Authors	Year	Number	Statin	Number of	Inductor	Start of statin	Duration	Evaluations	Results
		of		times per			$\mathbf{0f}$		
		animals		day			treatment		
Lee	2008	28	Control=9	Once a day	KA (10 mg/kg)	Before KA	7 days	Racine scale	Treatment with ATV efficiently
<i>et al</i> . ^[11]			ATV 10 mg/kg			injection- ATV			decreased convulsive events
			pre-SE=11			pre-SE group;			induced by KA, the neuronal death
			ATV 10 mg/kg			0.5 h after KA			of the hippocampus, infiltration of
			Post-SE=8			in ATV Post-SE			monocytes and proinflammatory
						group			gene expression
Gouveia	2011	20	Control=5, Solo	2 h after	Pilocarpine	After	0,5 days	Quantitative real-time PCR	LVT significant decrease in
<i>et al</i> . ^[9]			LVT 20 mg=5,	the onset of	(350 mg/kg)- 1	pilocarpine		(for cytokines and kinin	mRNA expression of IL-1 β ,
			Solo pilocarpine=5,	SE and 12	mg/kg before	injection		B1 and B2 receptors) and	IL-6, TNF- α , and kinin B1
			pilocarpine plus	h after the	scopolamine			corporal temperature	receptor, also reduced SE-induced
			LVT=5	first dose	methyl nitrate				hyperthermia
Campos	2017	152	Control=14, ATV	Once a day	Pilocarpine	After	14 days	Giemsa staining	The ATV regarded against
<i>et al</i> . ^[19]			10 mg/kg=10, ATV		100 mg/kg and	pilocarpine			tonic-clonic seizures caused by
			100 mg/kg=10,		Pilocarpine (30	injection			PTZ on the 14th post-SE. The
			Solo SE=13, ATV		mg/kg)				protective effects were similar in
			10 + SE = 18, ATV						female and male mice, requiring a
			100 mg/kg + SE = 15						higher amount of ATV in females
									(100 mg/kg versus 10 mg/kg in
									males)
PTZ: Pentyl Polymerase	lenetetra chain re	tzole, ICES: saction, TN	Increasing current elect F- α : Tumor necrosis fa	troshock, ATV: ctor alpha, IL-	: Atorvastatin, ES: Elε 1β: Interleukin-1β, IN	sctrical stimulation, E $VF \gamma$: Interferon gam	.EG: Electro na, GABA:	sncephalography, SE: Status ep gamma-Aminobutyric acid, K	oilepticus, LVT: Levetiracetam, PCR: A: Kainic acid, QA: Quinolinic acid
					-				

		Table 2: P	ositive	results rep	orted in experiment	al studies	
Study	Neuro- inflammation	Seizure prevention	Cell death	Prevention of kindling	Increase in convulsive threshold	Increase in latency and decrease in frequency of crisis	Reduction in mortality
Sehar et al. ^[1]				•	•	× •	
Oliveira et al.[3]	•						
Akgün et al. ^[2]	•			•		•	•
Gouveia et al. ^[9]	•						
Piermartiri et al.[10]		•	•				
Lee <i>et al</i> . ^[11]	•		•				
Gouveia et al.[5]	•						
Shafaroodi et al.[12]					•		•
Seker et al. ^[13]	•						
Moezi et al.[21]					•		•

•The study evaluated the respective parameter, finding positives results after the statin administration

			Ta	ble 3: Statin stu	idies in hu	mans		
Authors	Year	Type of study	Period evaluated	Number of patients	Age	Statin	Objective	Results
Pugh et al. ^[14]	2009	Retrospective cohort study	October 1999-September 2000	Cohort of epilepsy (n=1847) cohort without epilepsy (n=1,023,376)	≥65	Unspecified	Identify risk factors for new-onset geriatric epilepsy	The prescription of statins showed an OR=0.64, 95% CI=0.56-0.73 for the prevention of epilepsy
Etminan et al. ^[15]	2010	Nested case-control study	1995-2004	217 cases and 2117 controls	69.4±12.3 (cases) 70.0±9.6 (controls)	Atorvastatin, lovastatin, cerivastatin, fluvastatin, pravastatin, simvastatin, and rosuvastatin	To assess the potential efficacy of statins in the prevention of epilepsy	The ARR in patients with current use of statins was 0.65 with 95% CI 0.46-0.92, The ARR for previous users of statins was 0.72 (95% CI 0.39-1.30)
Sierra-Marcos et al. ^[16]	2014	Retrospective cohort study	April 2006-September 2012	427 patients	60.9±17.8	Simvastatin, atorvastatin, and pravastatin	To evaluate the possible role modulator of statins in status epilepticus	Statins were associated with lower mortality (relative risk ratio 0.38 , P=0.046)
Trivedi et al. ^[17]	2018	Retrospective cohort study	October 2003-March 2012	43,438 patients	56.0±12.0 statin users 55.7±12.4 No statin users	Simvastatin, atorvastatin, pravastatin, and rosuvastatin	To examine the association between the use of statins and the risk of epilepsy	The OR of epilepsy in the general cohort was 0.91 (95% CI=0.67-1.23) and in the healthy cohort, it was 1.08 (95% CI=0.64-1, 83). No significant beneficial or detrimental effect of the use of statins on the risk of epilepsy was demonstrated

ARR: Adjusted rate ratio, OR: Odds ratio, CI: Confidence interval

studies used ATV with a difference in the duration of treatment. However, both studies found a reduction in IL-1 β and TNF- α .^[5,9]

Statins and nitric oxide

Four studies evaluated the effect of statins on nitric oxide. Three studies used PTZ as an inducer and one used penicillin G ATV.^[13,16,17,20] Akgün Dar *et al*,

administered ATV 30 min before induction found that pretreatment with this statin reduced the inducible nitric oxide synthetase and matrix metalloproteinase 2 that have been related to a proconvulsant activity (P < 0.001) and also found a reduction in frequency and an increase in latency of seizures (P < 0.001).^[2] These findings agree with the results of the study by Shafaroodi *et al.*, who used PTZ and electrical stimulation as inductors and

Route of administration



Figure 6: Statin route of administration

administered ATV of 10 and 20 mg/kg; this treatment increased the threshold for tonic seizures caused by PTZ and decreased the appearance of tonic seizures and death in the induction model with electric current.^[12] Similarly, the study by Moezi et al. found an increase in the seizure threshold for both intravenous and intraperitoneal PTZ models and a decrease in the appearance of tonic seizures and death in the chronically treated group with ATV and induction with intraperitoneal PTZ.^[21] In the study by Seker et al., ATV, simvastatin, and rosuvastatin were used at a dose of 20 mg/kg; in this case, the group with rosuvastatin presented the best antiepileptic effect with a decrease in the expression of p53, Bax, and caspase 3 that are associated with cellular apoptosis, together with an increase in endothelial nitric oxide synthetase.^[13] Three studies (Seker et al., Shafaroodi et al., and Moezi et al.) showed that inhibitors of nitric oxide synthetase (L-NAME and aminoguanidine) decreased the anticonvulsant effect of ATV.^[12,13,21]

Statins and specific syndromes associated with epilepsy

We identified two studies that evaluated the use of statins in genetic syndromes with epileptogenic characteristics. In 2013, Osterweil *et al.* found that the administration of 100 mg/kg of lovastatin to genetically modified mice (fragile X syndrome) was able to correct the excessive synthesis of proteins and prevent the outbreak of epileptiform activity in the hippocampus *in vitro*, in addition to protecting the mice *in vivo*.^[1,22] On the other hand, in 2018, it was postulated that lovastatin is capable of modulating upregulated protein functions in animal models with Angelman syndrome by a mechanism other than the inhibition of protein synthesis.^[23] However, more research is needed to clarify this relationship.

Statin dose used in studies with animal models



MG

MG

MG

DISCUSSION

The neuroprotective effects of statins have been observed in various neurological diseases. However, its protective effect on epilepsy continues to be debated and most studies have been conducted in rats. An association has been found between the use of statins and the reduction of the risk of epilepsy in older adults. Sierra-Marcos *et al.* demonstrated an antiepileptic role of statins in older adults, for which they used the registry of 427 patients with an epileptic episode in a period of 6 years and took into account different predictive items of prognosis, among which if they had used or they use statins. The use of statins was statistically significant, which correlated with a decrease in morbidity and mortality in these patients.^[16]

Etminan *et al.* found that statins reduce the risk of hospitalization in patients with epilepsy focused on senile patients and stipulated that the result is directly related to the dose and the possible anti-inflammatory properties that statins confer.^[15] The few studies conducted in humans focused on observing the properties of statins in terms of epilepsy have had an effect in elderly people who have been shown to have a certain risk of developing epilepsy associated with cerebrovascular diseases and dementia. Pugh *et al.*, in their study on the risk factors of epilepsy of onset in old age, observed that in patients with prescription of statins, the development of epilepsies was lower.^[14]

There is research in humans that suggests the neuroprotective benefit of statins that is generalized to the entire population. Trivedi *et al.* could not determine the benefit of statins in their study where they included healthy individuals or those with few comorbidities and who used statins or not; however, it is emphasized that they were not associated with an increase in the risk of statins. Development of epilepsies which allows establishing that the use of statins is safe in patients who have epilepsy.^[17] All studies reviewed in humans agree on the need for more research on the subject to be able to define the beneficial effect of statins and be able to explain it.

Summary of evidence

This systematic review gives an overview of the available literature on the role of statins and its possible effect on epilepsy. This study only provides evidence 3b.

Limitations

Our study has some limitations. Most studies focused largely on an experimental level. All articles included in this review are peer-reviewed. There is a possibility of publication bias. Finally, the inclusion of only articles in English and Spanish could affect the generalization of our findings.

CONCLUSIONS

The role as anticonvulsant agents of statins is not completely known. Still, there is missing evidence to know how this type of mechanism works modulating the immune system. The participation of statins as reducing agents of neuroinflammation requires more studies.

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Conflicts of interest

There are no conflicts of interest.

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