



Original Article

Association between chronic stress and immune response to influenza vaccine in healthcare workers



Ema Sacadura-Leite^{a,b,c,*}, António Sousa-Uva^{b,c}, Helena Rebelo-de-Andrade^{d,e}, Sancha Ferreira^a, Regina Rocha^a

^a Occupational Health Department of Hospital de Santa Maria/CHLN, Lisbon, Portugal

^b National School of Public Health/New University of Lisbon, Lisbon, Portugal

^c CMDT – Centro de Investigação da Malária e Doenças Tropicais – Public Health, Lisbon, Portugal

^d National Institute of Health, INSA, Lisbon, Portugal

^e Faculty of Pharmacy, University of Lisbon, Lisbon, Portugal

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ABSTRACT

Introduction: Chronic stress can influence immune response to vaccines. Healthcare workers are exposed to stressors and biological hazards, the health effects of which may be prevented through vaccination.

Objectives: This study aims to evaluate the association between stress in nurses and: (1) insufficient response to influenza vaccine, assessed one month after vaccination (T_1); (2) the drop in haemagglutination-inhibition (HAI) antibodies (ab) six months after vaccination (T_6).

Methods: A nested case-control study was carried out with 136 healthy hospital nurses. Individual interviews, the *General Health Questionnaire* (GHQ₁₂) and *Maslach Burnout Inventory* (MBI-HSS) were applied in order to determine the presence of stress, using the triangulation method at the beginning of the study (T_0). Influenza vaccine was administered and titres of HAI above each strain composing influenza vaccine before vaccination (T_0), at T_1 and T_6 were assessed.

Results: There was no statistically relevant (5%) relationship between stress and the insufficient immune response to the vaccine at T_1 . Nevertheless, there was an association between stress and the drop in HAI ab AH₁ at T_6 , when we assessed stress by the triangulation method using an interview ($p = 0.006$), GHQ₁₂ ($p = 0.045$) and combination of criteria ($p = 0.001$), even after multivariate analysis (respectively, $p = 0.01$, $p < 0.05$ and $p = 0.002$). The odds ratios were, respectively, 3.64, 2.73 and 5.22.

Conclusions: The association we found, between chronic stress and the drop in HAI ab at T_6 , corroborates the hypothesis that stress can negatively influence immune response. Therefore, it seems reasonable to consider this issue when we implement vaccination programmes for healthcare workers.

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* Corresponding author.

E-mail address: ema.leite@chln.min-saude.pt (E. Sacadura-Leite).

Stresse crónico e a imunidade à vacina contra a gripe em profissionais de saúde

R E S U M O

Palavras-chave:

Anticorpos
Profissionais de saúde
Vacina contra a gripe
Stresse

Introdução: O stresse crónico pode influenciar a resposta imunitária à vacinação. Os profissionais de saúde estão expostos a stressores de natureza profissional e ainda a agentes biológicos cujos efeitos poderão ser prevenidos pela vacinação.

Objetivos: Estudar a associação entre a presença de stresse e (1) a “insuficiente” resposta imunitária à vacina contra a gripe, avaliada um mês após a vacinação (T_1); (2) a redução dos títulos de anticorpos dirigidos às hemaglutininas (HAI) seis meses após a vacinação (T_6).

Métodos: Realizou-se um estudo caso-controlo incorporado num estudo de coortes com a participação de 136 enfermeiros hospitalares saudáveis. Realizaram-se entrevistas individuais e aplicaram-se os questionários *The General Health Questionnaire* (GHQ₁₂) e *Maslach Burnout Inventory* (MBI-HSS) para determinação da presença de stresse crónico pelo método da triangulação, no início do estudo (T_0). Foi administrada a vacina contra a gripe e determinou-se os títulos de HAI dirigidos a cada estirpe componentes da vacina contra a gripe, antes da vacinação (T_0), em T_1 e em T_6 .

Resultados: Não se encontrou associação significativa (5%) entre a presença de stress e a “insuficiente” resposta à vacina contra a gripe em T_1 . Contudo, encontrou-se uma associação entre a presença de stress e a diminuição do título de HAI dirigidos à estirpe A(H₁N₁) em T_6 quando se avaliou a presença de stresse pelo método da triangulação usando a entrevista ($p=0,006$), o GHQ₁₂ ($p=0,045$) e a combinação dos três critérios ($p=0,001$), que se manteve após análise multivariada (respetivamente $p=0,01$, $p<0,05$ e $p=0,002$). Os odds ratio ajustados foram de 3,64, de 2,73 e de 5,22.

Conclusões: A associação encontrada entre a presença de stresse crónico e a redução do título de HAI em T_6 vem apoiar a hipótese de que o stresse poderá influenciar negativamente a manutenção dos títulos de anticorpos, mesmo em indivíduos adultos não idosos. Assim, parece razoável considerar este aspeto quando se pretende implementar programas de vacinação dirigidos a profissionais de saúde.

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Introduction

Healthcare workers are exposed to many stressors, some of them related with organisational work conditions and others, more specific to this profession, related with their activity of caring for the ill.¹⁻³

Chronic stress and burnout seem to be very common in nurses.⁴⁻⁶ For example, López-Castillo and colleagues found high levels of emotional disturbance determined by the General Health Questionnaire (GHQ₂₈) in 39% of hospital nurses.⁷

Amongst the consequences of chronic distress, whether they are related or not with work, are the possible effects on the immune system, including effects on the immune response to vaccination.

Healthcare workers are strongly advised to be vaccinated against influenza in order to protect themselves against the disease, reduce staff absenteeism and minimise the risk of nosocomial transmission to the patients they take care of. Vaccination is a possible model for immune response, testing mostly the humoral immune response. Vaccinated people develop antibodies (ab) that bind and neutralise the virus, in most cases ab against the surface glycoprotein hemagglutinin. Those ab can be used as markers of protection against the disease,⁸ caused by strains that are similar to the vaccine composition.

According to meta-analysis by Segerstrom and Miller, chronic exposure to stressors such as taking care of spouses with dementia, unemployment and suffering from physical disability is associated with a smaller ab response to influenza vaccine.⁹ Some reviews also suggest that chronic stress is associated with a smaller ab response to influenza vaccine.¹⁰⁻¹²

Generally speaking, studies evaluating the association between chronic stress and immune response to influenza vaccine showed relatively consistent results in old people. In those people, chronic exposure to stressors was associated with chronic anxiety and symptoms of depression and a lower response to influenza vaccine, in comparison to a control group.¹³⁻¹⁷

The use of a standardized dose of antigen which promotes a good immune response in most adults, could make it difficult the detection of the influence of chronic stress in the immune response to vaccination in younger adults, with a robust immune system.

Older people have a weaker immune system, related with age, so this could be an explanation for the greater consistency of results showing a negative association between chronic stress and immune response to vaccines in them. Vedhara and colleagues did not find any association between taking care of spouses with multiple sclerosis and ab response to influenza vaccine in adults under the age of 55.¹⁸

However, those adults showed similar stress levels as the control group.

In younger adults, such as university students, some studies¹⁹⁻²² found an association between stress, characterised in different ways, and the immune response to influenza vaccine (assessed by ab titres or by response rate) one month after vaccination. However, other studies did not find that association.²³⁻²⁵

Some of the studies found an association between stress and a drop in ab titres assessed 4-6 months after vaccination,^{19-23,25} even in the youngest adults. The drop in the ab titres associated with stress was only observed against one strain of vaccine components, suggesting that different exposures or different past vaccinations can be responsible for those results.

In an occupational context of healthcare settings, where healthcare workers are in the labour market (and are therefore not very old), but are simultaneously exposed to chronic stressors and biological risk hazards, it seems important to study the influence of stress on immune response to influenza vaccine.

Therefore, this study analyses the association between stress in nurses and: (1) insufficient response to influenza vaccine, assessed one month after vaccination (T_1); (2) the drop in influenza haemagglutination-inhibition (HAI) ab titres six months after vaccination (T_6), as compared to one month post vaccination HAI ab titres.

Materials and methods

Study design and participants

The study was a nested case-control study, conducted over six months in a university hospital during the 2007/2008 influenza season. Subjects were hospital nurses who were not taking any regular medication, including drugs that could affect immunity (such as cancer therapy drugs or corticosteroids). They did not have any medical condition that could affect the immune response and they also had no major surgery in the preceding three months. They did not have a history of drug consumption or alcohol consumption greater than 10 units per week, nor did they handle cytotoxic drugs or work regularly with ionising radiation ($n = 136$). The hospital's Ethics Committee approved the study and all the participants signed their agreement to participating in it.

One-month and six-month drop-out criteria:

- clinical diagnosis of medical condition that may affect immune response after the beginning of the study or taking any regular medication that can affect immunity (assessed by interview, at T_1 and T_6);
- workplace changes with regular exposure to ionising radiations or cytotoxic drugs handling;
- clinical influenza symptoms with virus identification in nasal or oropharyngeal swab, during the six months of the study;
- a rise in HAI Ab titre to A(H_1N_1), A(H_3N_2) or B strains six months after vaccination, as compared with the titres measured one month after vaccination. Such a rise in HAI Ab

titre suggests an exposure to influenza strains during T_1 and T_6 instead of a delayed response to the vaccine.

Stress assessment

Structured individual interviews were conducted at the beginning of the study (T_0) in order to:

- identify socio-demographic characteristics and possible confoundable variables related with immunity (physical exercise, nutritional parameters, nutritional supplements, hours of sleep per day, smoking habits, shift work) and influenza vaccination history;
- identify work-related and non-work-related stressors, using a Likert scale from 1 to 5;
- assess perceived stress, using a Likert scale from 1 to 5;
- identify stress-related behavioural changes or psychosomatic symptoms.

We also applied the Portuguese versions of the *General Health Questionnaire* (GHQ₁₂) and *Maslach Burnout Inventory* (MBI-HSS or MBI) exhaustion scale at the beginning of the study (T_0). Alfa Cronbach for those scales was 0.855 and 0.874 respectively.

In order to assess the presence of chronic stress, we applied the triangulation method at T_0 , as suggested by Cox and colleagues,²⁶ in four different ways:

- through interviews: we accepted the presence of chronic stress using interviews if there were identified stressors classified as 4 or 5, plus perceived stress classified as 4 or 5, plus at least one behavioural change or one psychosomatic symptom stress-related;
- through GHQ₁₂: we accepted the presence of chronic stress using GHQ₁₂ if there were identified stressors classified as 4 or 5, plus GHQ₁₂ higher than 2, plus at least one behavioural change or one psychosomatic symptom stress-related;
- through MBI: we accepted the presence of chronic stress using MBI if there were identified stressors classified as 4 or 5, plus MBI exhaustion scale higher than 24, plus at least one behavioural change or one psychosomatic symptom stress-related;
- combination of criteria: we accepted the presence of stress using combination of criteria if there was stress using interview and stress using GHQ₁₂ or if there was stress using interview and stress using MBI exhaustion scale.

Vaccination and laboratory procedures

Venous blood was drawn at three stages between October 2007 and April 2008: (i) immediately before influenza vaccination (T_0); (ii) one month following immunisation (T_1); and (iii) six months after T_0 (T_6).

The samples rested 1 h at ambient temperature following centrifugation at 3500 rpm for 10 min. Sera were stored at -20°C until used. All the samples drawn at T_0 , T_1 and T_6 were processed at the same time and under the same conditions.

A commercially available 2007/2008 trivalent influenza vaccine, with the recommended composition for that season in North Hemisphere, was administered intramuscularly, in the

deltoid muscle, during October. All the vaccines belonged to the same group (AFLUA290AD).

Haemagglutination inhibition reaction was used to assess specific HAI ab titre against influenza A(H₁N₁), A(H₃N₂) and B strains included in the influenza vaccine, in accordance with the World Health Organisation's manual.²⁷ Immediately before the laboratorial procedures, the sera were treated by a Receptor-Destroying Enzyme (RDE) in order to remove unspecific agglutinins and inhibitors.

The reference antigens were diluted to obtain 4 units against haemmagglutinin per 25 µl and incubated with the treated sera samples. Erythrocytes were then added to the fluid.

HAI Ab titre corresponded to the inverse of the last dilution of serum that completely inhibited haemagglutination. We used progressive dilutions, starting with 1:10 up to 1:20.480.

The serological parameters obtained were:

- HAI Ab titre against influenza A(H₁N₁), A(H₃N₂) and B strains included in influenza vaccine, before (T₀) and after vaccination (T₁ and T₆);
- rise in HAI Ab titre against influenza A(H₁N₁), A(H₃N₂) and B strains included in influenza vaccine, assessed one month after vaccination (compared to HAI Ab titre immediately before vaccination);
- drop in HAI Ab titre against influenza A(H₁N₁), A(H₃N₂) and B strains included in influenza vaccine, between T₁ and T₆.

Definitions

For analysis at T₁ and at T₆, we defined the following groups:

responders at T₁ (one month after vaccination): participants that showed, at T₁, at least a fourfold rise in HAI ab titre compared to the titre before vaccination;

non-responders at T₁ (one month after vaccination): participants that did not show, at T₁, a fourfold rise in HAI ab titre compared to the titre before vaccination;

HAI ab titre drop group at T₆ (six months after vaccination): participants with at least a fourfold rise in HAI ab titre at T₁, but who showed a drop in HAI ab titre at T₆, as compared to HAI ab titre at T₁;

no change in HAI ab titre group at T₆ (six months after vaccination): participants with at least a fourfold rise in HAI ab titre at T₁, but with no change in HAI ab titre at T₆, as compared to HAI ab titre at T₁.

Statistical analyses

For dichotomous variables, we used the *Qui-square* and Fisher exact tests and determined the *odds ratio*.

For numerical variables, we used the *Kolmogorov-Smirnov* and *Shapiro-Wilk* tests to assess normal distribution in non-responders and responders (T₁) and in the HAI Ab titre drop group and no change in HAI Ab titre group (T₆).

For normal distributions, the *T Student* test was used to compare means, and the *Levene* test to assess variance homogeneity. For no normal distributions, we applied the *Mann-Whitney* test to compare medians.

We also used the multivariate analysis and determined the adjusted odds ratio for the confounding variables.

We considered a statistical significance of 5%. All tests were run in the *Statistical Package for Social Sciences – SPSS®* software, version 14.0 for Windows.

Results

We studied 136 nurses, most of whom were female (83.8%), Caucasian (96.3%) and did not smoke (77.9%). Their average age was 33 and the median age was 29 (22–63 years old). Only one participant was more than sixty years of age.

More than one half of the participants had been given an influenza vaccine shot at least one of the four seasons prior to the beginning of the study (52.9%), mostly in the year immediately before (44.1%). Nurses included in the study worked mostly on a shift work basis (70.6%), had a corporal index mass (kg/m²) between 18.5 and 24.9 (72.1%) and slept at least 7 h per day (66.2%). The majority did not take vitamin supplements (86.8%) or fish oil (98.5%). Only 54.4% did regular physical exercise and 46.3% ate yogurt daily.

Association between chronic stress and non-responders at T₁

One month after vaccination (T₁), we did not find any association of statistical significance between non-responders for A(H₁N₁) virus strain included in the influenza vaccine and the presence of chronic stress at T₀, assessed in four different ways. Similarly, there was also no association between non-responders for A(H₃N₂) or non-responders for B strains included in the influenza vaccine and chronic stress at T₀. To simplify the table, we named responders or non-responders for A(H₁N₁) and for A(H₃N₂) as responders or non-responders AH₁ and AH₃ respectively (Table 1).

Aemagglutination-inhibition antibodies titres at T₀ in responders and non-responders at T₁

Using the *Kolmogorov-Smirnov* and *Shapiro-Wilk* tests, we found that there is no normal distribution in non-responders and responders at T₁ for the considered continuous variables (*p* < 0.001). Therefore, we applied the *Mann-Whitney* test to compare medians between HAI ab titres at T₀ in responders and non-responders at T₁. We found that non-responders AH₁ at T₁ had significantly higher HAI ab AH₁ titres at T₀ than responders AH₁ at T₁. The same happened with non-responders AH₃ at T₁ and non-responders B at T₁, who had significantly higher HAI AH₃ titres at T₀ and HAI B titres at T₀ than the corresponding responders (Table 2).

Association between chronic stress and drop in aemagglutination-inhibition ab titres at T₆

At T₆ (six months after vaccination), the presence of stress in the HAI ab AH₁ titre drop group was higher than in the no change in HAI ab AH₁ titre group, when we assessed stress by all the considered different ways, being statistically significant when we assessed the presence of chronic stress by

Table 1 – Stress in non-responders and in responders AH₁, AH₃ and B at T₁.

HAI ab groups at T ₁	Chronic stress assessment	Stress in non-responders at T ₁	Stress in responders at T ₁	Statistical analysis (Qui-square test)	
		n (%)	n (%)	OR (95% CI)	p
AH ₁ (n = 135)	Stress (interview)	28 (62.2)	46 (51.1)	1.575 (0.759–3.272)	0.221
Non-responders: 45	Stress (GHQ ₁₂) ^a	19 (42.2)	44 (48.9)	0.764 (0.371–1.572)	0.464
Responders: 90	Stress (MBI) ^b	16 (35.6)	36 (40.0)	0.828 (0.394–1.738)	0.617
	Stress (combination of criteria) ^c	20 (44.4)	41 (45.6)	0.956 (0.466–1.963)	0.903
AH ₃ (n = 136)	Stress (interview)	27 (54.0)	47 (54.7)	0.974 (0.484–1.961)	0.941
Non-responders: 50	Stress (GHQ ₁₂) ^a	20 (40.0)	43 (50.0)	0.667 (0.329–1.351)	0.259
Responders: 86	Stress (MBI) ^b	15 (30.0)	37 (43.0)	0.568 (0.271–1.190)	0.132
	Stress (combination criteria) ^c	20 (40.0)	41 (47.7)	0.732 (0.361–1.483)	0.386
B (n = 135)	Stress (interview)	31 (52.5)	43 (56.6)	0.850 (0.429–1.683)	0.640
Non-responders: 59	Stress (GHQ ₁₂) ^a	24 (40.7)	39 (51.3)	0.651 (0.327–1.293)	0.219
Responders: 76	Stress (MBI) ^b	22 (37.3)	30 (39.5)	0.912 (0.453–1.836)	0.796
	Stress (combination of criteria) ^c	24 (40.7)	37 (48.7)	0.723 (0.364–1.437)	0.354

OR – odds ratio; 95% CI – 95% confidence interval.

^a General Health Questionnaire (GHQ₁₂).

^b Maslach Burnout Inventory (MBI).

^c Stress (interview and GHQ₁₂) or stress (interview and MBI).

triangulation method at T₀ using interviews, GHQ₁₂ and the combination of the three methods. On the contrary, we did not find any statistically significant association between the others HAI ab titre drop groups at T₆ and the presence of chronic stress (Table 3).

Association between other possible confounding variables and aemagglutination-inhibition ab AH₁ titre drop group at T₆

Some conditions that can affect immunity could have been possible confounding factors, when we considered the association between stress and the HAI ab AH₁ titre drop group at T₆. Using the Kolmogorov–Smirnov and Shapiro–Wilk tests, we found that there is no normal distribution in HAI ab AH₁ titre drop group at T₆ and in no change in HAI ab AH₁ titre at T₆ for the considered continuous variables ($p < 0.001$). Therefore, we applied the Mann–Whitney test to compare their medians.

The statistical analyses did not find any significant difference between groups at T₆ for the variables (continuous and dichotomous) taken into consideration (Table 4).

Multivariate analysis and adjusted odds ratios for stress, age and aemagglutination-inhibition ab AH₁ titres at T₀ and T₁ considering aemagglutination-inhibition ab AH₁ titre drop group at T₆

Stress – assessed by interview, GHQ₁₂ or using the combination of the three methods – was the exclusive variable associated with HAI ab AH₁ titre drop group at T₆, but we also took the age variable into consideration in the multivariate analysis. That option was made because age is a strong factor influencing immunity and, in the simple analysis, the median difference between groups would be different if we considered a confidence level of 90% (instead of 95%).

Furthermore, Beyer and colleagues showed that basal HAI ab AH₁ titres influence HAI ab AH₁ titres one month after

Table 2 – HAI antibodies AH₁, AH₃ and B titres at T₀ in non-responders and in responders AH₁, AH₃ and B at T₁.

HAI antibodies at T ₀	Non-responders at T ₁	Responders at T ₁	Statistical analysis (Mann–Whitney test)	
	Md (min–max)	Md (min–max)	Md differences	p
HAI ab AH ₁ titres at T ₀ in non-responders AH ₁ (n = 45) and responders AH ₁ (n = 90) at T ₁	640.0 (80–10,240)	20.0 (10–1280)	620.0	<0001
HAI ab AH ₃ titres at T ₀ in non-responders AH ₃ (n = 50) and responders AH ₃ (n = 86) at T ₁	160.0 (10–5120)	20.0 (10–1280)	140.0	<0001
HAI ab B titres at T ₀ in non-responders B (n = 59) and responders B (n = 76) at T ₁	160.0 (20–10,240)	60.0 (10–640)	100.0	<0001

Md - median.

Table 3 – Stress in HAI antibodies titre drop group and in no change in HAI Ab titre group AH₁, AH₃ and B at T₆.

HAI ab groups at T ₆	Chronic stress assessment	Stress in HAI ab titre drop group at T ₆	Stress in no change in HAI Ab titre group at T ₆	Statistical analysis (Qui-square test)	
		n (%)	n (%)	OR (95% CI)	p
AH ₁ (n = 88)	Stress (interview)	36 (63.2)	10 (32.3)	3.600 (1.427–9.084)	0.006
HAI ab titre drop group: 57	Stress (GHQ ₁₂) ^a	33 (57.9)	11 (35.5)	2.500 (1.012–6.176)	0.045
	Stress (MBI) ^b	25 (43.9)	10 (32.3)	1.641 (0.656–4.104)	0.288
No change in HAI ab titre group: 31	Stress (combination of criteria) ^c	34 (59.6)	7 (22.6)	5.068 (1.875–13.70)	0.001
AH ₃ (n = 81)	Stress (interview)	25 (52.1)	21 (63.6)	0.621 (0.251–1.539)	0.302
HAI ab titre drop group: 48	Stress (GHQ ₁₂) ^a	26 (54.2)	(51.5)	1.112 (0.458–2.703)	0.814
	Stress (MBI) ^b	20 (41.7)	16 (48.5)	0.759 (0.311–1.851)	0.544
No change in HAI ab titre group: 33	Stress (combination of criteria) ^c	23 (47.9)	18 (54.5)	0.767 (0.315–1.865)	0.558
B (n = 76)	Stress (interview)	26 (51.0)	17 (68.0)	0.489 (0.179–1.335)	0.160
HAI ab titre drop group: 51	Stress (GHQ ₁₂) ^a	27 (52.9)	12 (48.0)	1.219 (0.468–3.177)	0.686
	Stress (MBI) ^b	17 (33.3)	13 (52.0)	0.462 (0.174–1.226)	0.118
No change in HAI ab titre group: 25	Stress (combination of criteria) ^c	22 (43.1)	15 (60.0)	0.506 (0.191–1.339)	0.167

OR – odds ratio; 95% CI – 95% confidence interval.

^a General Health Questionnaire (GHQ₁₂).

^b Maslach Burnout Inventory (MBI).

^c Stress (interview and GHQ₁₂) or stress (interview and MBI).

Table 4 – Variables distribution (stress not included) in HAI ab AH₁ titre drop group at T₆ and in no change in HAI ab AH₁ titre group AH₁ at T₆.

Variables	HAI ab AH ₁ titre drop group at T ₆ (n = 57)	No change in HAI ab AH ₁ titre group at T ₆ (n = 31)	Statistical analysis (Fisher test or Qui-square test or Mann–Whitney test)	
	n (%)	n (%)	OR (95% CI)	p
Male gender	5 (8.8)	7 (22.6)	0.330 (0.095–1.145)	0.103 ^a
Caucasian	53 (93.0)	31 (100)	0.631 (0.536–0.743)	0.293 ^a
Shift work	35 (61.4)	23 (74.2)	0.553 (0.211–1.453)	0.250 ^a
Daily sleep hours < 7	20 (35.1)	8 (25.8)	0.643 (0.244–1.699)	0.372 ^b
Smokers	10 (17.5)	9 (29.0)	0.520 (0.185–1.461)	0.211 ^b
Vitamin supplements	10 (17.5)	4 (12.9)	1.436 (0.410–5.025)	0.762 ^a
Fish oil consumption	0 (0.0)	0 (0.0)		
Daily consumption of yogurts	27 (47.4)	15 (48.4)	0.960 (0.400–2.304)	0.927 ^b
No regular physical activity	29 (50.9)	11 (35.5)	1.883 (0.765–4.634)	0.166 ^b
Past influenza vaccine	17 (29.8)	12 (38.7)	0.673 (0.268–1.687)	0.397 ^b
Influenza vaccine at 2006	14 (24.6)	11 (35.5)	0.592 (0.229–1.533)	0.278 ^b
HAI ab AH ₁ at T ₀ ≥ 40	25 (43.9)	17 (54.8)	0.643 (0.267–1.551)	0.325 ^b
Variables	HAI ab AH ₁ titre drop group at T ₆ (n = 57)	No change in HAI ab AH ₁ titre group at T ₆ (n = 31)	Statistical analysis (Fisher test or Qui-square test or Mann–Whitney test)	
	Md (min–max)	Md (min–max)	Md differences	p
Age	31.0 (23.0–63.0)	26.0 (22.0–56.0)	5.0	0.072 ^c
Body index mass	22.7 (17.7–37.8)	22.0 (18.0–37.5)	0.7	0.793 ^c
HAI ab AH ₁ titres at T ₀	20.0 (10–640)	40.0 (10–1280)	–20.0	0.276 ^c
HAI ab AH ₁ titres at T ₁	1280.0 (40–20,480)	1280.0 (160–20,480)	0.0	0.265 ^c

Md – median.

^a Fisher test.

^b Qui Square test.

^c Mann–Whitney test.

Table 5 – Multivaried analysis (multiple logistic regression) for stress (assessed by triangulation method using GHQ₁₂, using interview and using the combination of the three methods) in HAI ab AH₁ titre drop group at T₆ considering age, HAI ab AH₁ titres at T₀ and at T₁.

Chronic stress assessment	Considered variables in multivaried analysis	Statistical analysis (multiple logistic regression)	
		Adjusted OR (95% CI)	p
GHQ ₁₂ ^a	Age	1.038 (0.989–1.089)	0.134
	Stress	2.733 (1.039–7.186)	0.042
	HAI ab AH ₁ at T ₀	0.998 (0.995–1.000)	0.100
	HAI ab AH ₁ at T ₁	1.000 (1.000–1.000)	0.793
Interview	Age	1.043 (0.994–1.094)	0.083
	Stress	3.643 (1.371–9.684)	0.010
	HAI ab AH ₁ at T ₀	0.999 (0.996–1.001)	0.236
	HAI ab AH ₁ at T ₁	1.000 (1.000–1.000)	0.987
Combination of criteria ^b	Age	1.044 (0.994–1.096)	0.087
	Stress	5.223 (1.828–14.924)	0.002
	HAI ab AH ₁ at T ₀	0.999 (0.996–1.001)	0.255
	HAI ab AH ₁ at T ₁	1.000 (1.000–1.000)	0.892

Adjusted OR – adjusted odds ratio; 95% CI – 95% confidence interval.
^a General Health Questionnaire (GHQ₁₂).
^b Stress (interview and GHQ₁₂) or stress (interview and MBI).

vaccination.²⁸ Therefore, we also considered HAI ab AH₁ titres at T₀ and T₁ in the multivariate analysis.

We found that stress, assessed by triangulation method using GHQ₁₂, using interview and using the combination of the three methods, maintained the association with HAI ab AH₁ titre drop group at T₆. The association between HAI ab AH₁ titre drop group at T₆ and the others variables did not reveal any statistical significance (Table 5).

When we assessed stress by triangulation method using GHQ₁₂, the model was statistically significant ($p < 0.029$) and suitable, given that null hypothesis was not rejected in the Hosmer–Lemeshow test ($p < 0.106$). The model showed a validity rate of 65.9.

When we assessed stress by triangulation method using interview, the model was statistically significant ($p < 0.009$) and suitable, given that null hypothesis was not rejected in the Hosmer–Lemeshow test ($p < 0.761$). The model showed a validity rate of 71.6.

When we assessed stress by triangulation method using the combination of the three methods, the model was statistically significant ($p < 0.002$) and suitable, given that null hypothesis was not rejected in the Hosmer–Lemeshow test ($p < 0.679$). The model showed a validity rate of 73.9.

Discussion and conclusions

When human beings are exposed to chronic stressors, they may respond to them with neuroendocrine changes that include the release of neuropeptides, monoamines and hormones. Most of those substances are able to change the immune cells behaviour.²⁹

Psychological chronic stress can change antibody (ab) production and kinetics after vaccination, in particular after the influenza vaccine is given to elderly people who take care of spouses with dementia.^{9–12}

Various studies of elderly people have found an association between exposure to a long-term stressor (such as dementia spouse caregiving) and a small proportion of them who reached at least ab HAI titres that were fourfold what they had before flu vaccination, assessed one month after vaccination.^{13–15} Bereavement and marriage seem to be associated with antibody response to influenza vaccination in the elderly as well.¹⁷

In our study, as in some other studies involving young adults,^{18,23–25} we did not find any association between the presence of chronic stress in nurses and the proportion of them that reach at least four times the ab HAI titre levels they had before flu vaccination. On the contrary, other studies have found an association among perceived distress,¹⁹ life events,²⁰ neuroticism²¹ and loneliness²² and the immune response to flu vaccination, assessed one month or five weeks after.

It is possible that methodological issues can explain discrepancies in results verified in studies with younger groups, such as: (1) different ways of characterising independent variables; (2) samples with differing demographic characteristics and dimensions; (3) differing histories of flu virus exposure.

With respect to the latter issue, our study found that non-responders had significantly higher ab HAI AH₁N₁, AH₃N₂ and B titres at T₀ than responder groups. Therefore, as postulated by Beyer and colleagues,²⁸ ab HAI titres at T₀, showed to be an important confounding variable when studying the relationship between stress and immune response to flu vaccine one month after vaccination and must be considered.

Nevertheless, we found an association between the presence of chronic stress (measured in three different ways) and a drop in ab HAI (AH₁N₁) at T₆. Other studies also found an association between distress,^{19,25} life events,²⁰ life events weighed with perceived stress,²³ neuroticism,²¹ or loneliness²² and a drop in ab HAI four to six months after vaccination. Those associations were found for at least one strain composing the flu vaccine.

Our study found a large proportion of nurses with chronic stress in the HAI ab AH₁ titre drop group at T₆, as compared to the no change in HAI Ab AH₁ titre group at T₆, when we measured stress by triangulation method, using interview or GHQ₁₂ to assess perceived stress, and using the combination of the three methods (as described in the methods section). We did not find any statistically significant association when we assessed the presence of stress by the triangulation method but using the MBI exhaustion scale to measure perceived stress. A possible explanation is the fact that the MBI exhaustion scale measures specifically work-related stress and the possible immunologic repercussions of chronic stress seem to be independent of the stress source.

As described in other studies,^{19-23,25} the association between stress and a drop in ab HAI at T₆ did not occur for all the vaccine strains components.

Strain novelty can be an important factor in that analysis, as argued by other authors.^{15,20} Pressman and colleagues, for example, only found an association between stress and a drop in HAI ab, four months after vaccination, for a strain that was not included in previous vaccinations the participants received.²⁵ In our study, the exclusive strain that was not included in flu vaccines in the three preceding years was the A(H₁N₁) strain.

In Portugal the predominant circulating strains with high flu activity since 1990 have been A(H₃N₂) and B. From 1990 to the beginning of the study, the A(H₁N₁) strain was only predominant in 2005, simultaneously with strain B, and 2005 was a year with very low flu activity.³⁰ We also know that A strains undergo more drift mutations than B strains,³¹ so this can contribute to their being a relative novelty for the participants' immune system.

Finally, in our study, the A(H₁N₁) influenza strain proved to be the most immunogenic one, showing a rise in the HAI Ab titre geometric mean of 11.1. A (H₃N₂) and B strains showed rises of 6.2 and 4.6 times, respectively, between T₀ and T₁. It is possible that the best immunogenicity observed for the A (H₁N₁) strain was related with the fact that some participants had had a primary infection with an A (H₁N₁) strain, so the response to A (H₁N₁) antigens have been more robust in them.³² That could be a factor that may influence the detection of the association between stress and drops in HAI ab after a period of time.

Given that our sample was not a randomised sample because it depended on nurses voluntarily being vaccinated and participating in the study, we analysed distribution differences for some variables in the HAI ab AH₁ titre drop group at T₆ and the no change in HAI Ab AH₁ titre group at T₆ that are not included in drop out criteria. As there are a lot of variables for which we do not yet know if they can influence immunity, we studied those that are most referred to in the relevant literature.³³

We did not find any differences in the distribution of the studied variables in the two groups at T₆. Nevertheless, we decided to include ab AH₁ titres at T₀, ab AH₁ titres at T₁ and age in multiple logistic regression. The reason for including the first two variables was the strong suggestion in literature that they can influence ab titres after vaccination (immune response),²⁸ even though we found no references to the influence they have on a drop in titres six months after

flu vaccination. Age is strongly related with immunity³³ but we did not find any difference in terms of age between the two groups considered at T₆ at the significance level we considered (5%). If we considered a significance level of 10% the result would be different. Hence, we also included age in the multivariate analysis.

After the multivariate analysis, we still found an association with statistical significance between the presence of chronic stress and the HAI ab AH₁ titre drop group at T₆, when we assessed stress in three different ways, all of them using the triangulation method, as suggested by Cox and colleagues, as a good way of measuring stress.²⁶ Therefore, the relationship that we found between chronic stress and a drop in HAI ab at T₆ supports the thesis that stress can negatively influence HAI ab titres some months after flu vaccination even in people in adults under the age of 60. As we could notice, this is the first study assessing the association between chronic stress and immune response to influenza vaccine in healthcare workers, who is an important target group for influenza vaccine. Therefore, in an occupational health environment, it is reasonable to consider the possible interference of chronic stress with ab titres when we implement vaccination programmes to prevent biological occupational risks.

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Conflicts of interest

The authors have no conflicts of interest to declare.

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