

Anti-DFS70 Autoantibodies in Japanese Women at Different Menstrual Stages

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ABSTRACT

Peripheral arthralgia is a common complaint of peri- and post-menopausal women. Although the pathophysiology is not completely understood, it is thought that menopausal arthralgia may be related to estrogen deficiency and depletion. Menopausal women referred to rheumatologists with these complaints along with high titer anti-nuclear antibody (ANA) can lead to patient anxiety and present a diagnostic dilemma for the clinician. We report two index cases of menopausal women who had a positive ANA by indirect immunofluorescence (IIF) attributed to anti-dense fine speckled 70 (DFS70) autoantibody (Aab) that had developed arthralgias prompting referral to our clinic. It is known that the presence of this Aab is paradoxically found more frequently in healthy individuals compared to those with a systemic autoimmune rheumatic disease (SARD). The lack of evidence for a SARD in these patients and a remarkable response to hormone replacement therapy (HRT) prompted a wider study of anti-DFS70 Aab in peri- and post-menopausal women. Hence, a retrospective analysis of 282 Japanese female patients referred to an outpatient rheumatology clinic for arthralgias was performed. The results indicate that the frequency of anti-DFS70 Aab in peri- and post-menopausal women with non-SARD is higher than in other peri- and post-menopausal women with SARD. Therefore, testing for anti-DFS70 Aab in this population may be helpful in distinguishing between SARD and non-SARD patients with positive ANA, who may in fact, have menopausal-related arthralgia and are responsive to HRT.

Keywords: Antinuclear antibody (ANA), Anti-DFS70/LEDGF, Estrogen deficiency hormone replacement therapy, Menopausal arthralgia

Abbreviations: AAB: Autoantibody; ACR: American College of Rheumatology; ANA: Anti-Nuclear Antibody; DFS: Dense Fine Speckled; ELISA: Enzyme-Linked Immunoassay; EULAR: European League Against Rheumatism; FM: Fibromyalgia; FSH: Follicle Stimulating Hormone; HIV: Human Immunodeficiency Virus; HRT: Hormone Replacement Therapy; ICAP: International Consensus on ANA Patterns; IIF: Indirect Immunofluorescence; HTLV 1: Human T-Cell Leukemia Virus 1; mg: milligram; NASH: Non-Alcoholic Steatohepatitis; OD: Optical Density; OA: Osteoarthritis; PCNA: Proliferating Cell Nuclear Antigen; PeMA: Peri-Menopausal Arthralgia, PeMS: Peri-Menopausal Stage; PMS: Pre-Menopausal Syndrome; PoMA: Post-Menopausal Arthralgia, PoMS: Post-Menopausal Stage; PrMA: Pre-Menopausal Arthralgia, PrMS: Pre-Menopausal Stage; RA: Rheumatoid Arthritis; SARD: Systemic Autoimmune Disease; SjS: Sjögren's Syndrome; SLE: Systemic Lupus Erythematosus; SSc: Systemic Sclerosis; UA: Undifferentiated Arthralgia/Arthritis

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INTRODUCTION

Peripheral joint arthralgia along with menopausal symptoms of palpitations, hot flashes and diaphoresis are common complaints of peri- and post-menopausal women [1]. The cause of these musculoskeletal symptoms in this clinical setting is not completely established, although circumstantial evidence suggests it is associated with estrogen deficiency. In younger females, arthralgias often appear during the premenstrual stage as a part of premenstrual syndrome. Even in the peri-menstrual stage, menses become irregular because of fluctuating estrogen levels and the antioxidant α -tocopherol-N that is required to maintain regular periods [2]. In the post-menopausal stage, joint pains can be relieved by hormone replacement therapy (HRT) [3].

The clinical picture of arthralgia associated with menopause can become clinically challenging if a screening anti-nuclear antibody (ANA) by indirect immunofluorescence (IIF) test done to rule out a SARD is found to be positive. We present two index cases of menopausal females with arthralgia and high titer ANA. The ANA described in the sera of these two patients was characteristic of the dense fine speckled (DFS) pattern in nuclei of interphase cells and also of the metaphase plate of dividing cells (International Consensus on ANA Patterns IC-2: https://anapatterns.org/view_pattern.php?pattern=20) and on further testing the autoantibody (Aab) reactivity was confirmed to be directed against the 70 kDa DFS70 target autoantigen [4,5], also known as lens epithelium-derived growth factor (LEDGF) [5,6]. In general, this antibody is found in the sera of less than 10% of SARD patients but paradoxically found in 10-25% of healthy subjects [4,5,7,8]. On rheumatological evaluation, no evidence of SARD was found in these two patients and after two weeks of HRT, their arthralgias had completely resolved.

While anti-DFS70 Aab has been reported in wide range of diseases and healthy cohorts [4,5,7,8], no studies have focused on its presence in menopausal females. These two case reports of menopausal women with anti-DFS70 Aab without SARD prompted us to determine the frequency of anti-DFS70 Aab in pre-, peri- and post-menopausal females with and without SARD. Since the frequency of anti-DFS70 has been reported to decrease with age [9], but is higher in pre-pubertal females [10] a working hypothesis is that it should also have higher frequency in post-menopausal than in pre- or peri-menopausal females. Further, we set out to determine the frequency of anti-DFS70 Aab in females with undifferentiated arthritis (UA) as compared to patients with a defined SARD.

Patient 1

A 41 year old Japanese female was evaluated for recurrent chest discomfort and back pain. There was no evidence of pathology on chest X-ray or CT scan. Due to concerns of an underlying SARD, an ANA by IIF was done and was found

to be positive at a titer of 1:320 with a homogenous and speckled pattern. She had no other manifestations of a SARD. Six years later, her menopausal status was established and within 7 months she could not hold a pencil or pick up coins owing to finger and wrist joint pain. She was seen at a rheumatology clinic for an evaluation of possible rheumatoid arthritis (RA). The laboratory and clinical features are summarized in **Supplemental Figure 1a** which included a negative autoantibody profile except for a strongly positive anti-DFS70 Ab (OD index 147.1 units). A diagnosis of RA or other related-SARD was ruled out by the attending rheumatologist. After 2 months of HRT, her arthralgias completely resolved.

Patient 2

A 27 year old Japanese woman was noted to have a positive ANA by IIF at a titer of 1:160 with a homogenous and speckled pattern and after initial evaluation the diagnosis of SARD was ruled out. At the age of 52, her menses became irregular and within 5 months, she noticed joint pains localized to interphalangeal finger and bilateral knee joints. She was referred to a rheumatology clinic to evaluate the joint pains and laboratory and clinical assessment were unremarkable (**Supplemental Figure 1b**). In this peri-menopausal stage, she was treated with α -tocopherol N (600 mg/day) and non-steroidal anti-inflammatory medications which, within 2 months, gave her a 50% reduction of joint pain. However, as she entered full menopausal stage, she left her job as an office employee because of severe fatigue and anxiety. Within 2 weeks of cyclic HRT, she became asymptomatic and returned to her employment. Her follow-up ANA by IIF was positive at 1:80 with homogenous and speckled IIF pattern and further evaluation disclosed a positive anti-DFS70 antibody (OD index 80.9) test.

METHODS

Patient population

Two hundred and eighty-two female patients seen for evaluation of joint pain at the Keigu and Higashiterao Second Clinics (Yokohama, Japan) from November 2016 to October 2017 were enrolled and separated into 3 main clinical diagnostic groups: SARD, UA and non-SARD (e.g. Osteoarthritis (OA), Fibromyalgia (FM), healthy individuals (HI)) (**Figure 1**). A diagnosis of an SARD including RA [11], Sjögren's Syndrome (SjS) [12], Systemic Lupus Erythematosus (SLE) [13] or other SARDs was confirmed by a rheumatologist using appropriate classification criteria. Patients who did not meet full classification criteria for a SARD were classified as UA [14]. Patients with other non-SARD diseases such as OA, FM and healthy individuals were also included as controls. Ethical approval of this study was provided by The Society of Healthcare for Menopause and Aging in Tokyo, Japan and was compliant with the Helsinki Declaration of 1975 as revised in 2013:

<https://www.wma.net/policies-post/wma-declaration-of-helsinki-ethical-principles-for-medical-research-involving-human-subjects>

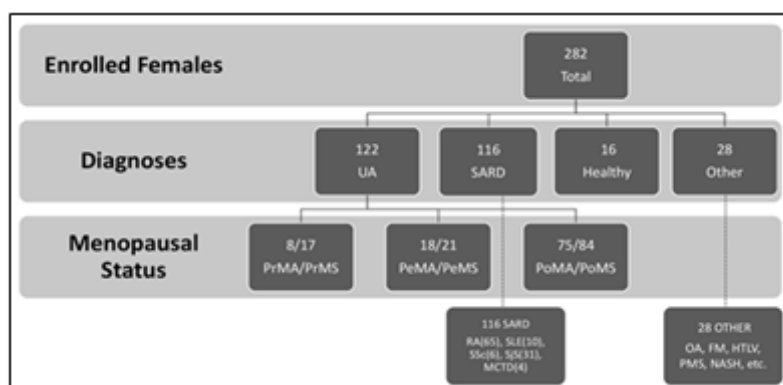


Figure 1. Distribution of 282 Japanese females enrolled in the anti-DFS70 study.

Abbreviations: FM: Fibromyalgia; HTLV 1: Human T-Cell Leukemia Virus 1; NASH: Non-Alcoholic Steatohepatitis; OA: Osteoarthritis; PeMA: Peri-Menopausal Arthralgia; PeMS: Peri-Menopausal Stage; PMS: Pre-Menopausal Syndrome; PoMA: Post-Menopausal Arthralgia, PoMS: Post-Menopausal Stage; PrMA: Pre-Menopausal Arthralgia; PrMS: Pre-Menopausal Stage; RA: Rheumatoid Arthritis; SARD: Systemic Autoimmune Disease; SjS: Sjögren's Syndrome; SLE: Systemic Lupus Erythematosus; SSc: Systemic Sclerosis; UA: Undifferentiated Arthralgia/Arthritis

Note: Normal healthy individuals and SARD patients had no menopausal symptoms at enrollment; HRT, tocopherol N and ultra-low dose pills were not prescribed. In case of less 30% reduction of joint pain through proper treatment, they remain UA at PrMS, PeMS and PoMS

Based on the regularity of menses, the UA patients were divided into three categories: Pre-Menopausal (PrM), Peri-Menopausal (PeM) and Post-Menopausal (PoM) stages. These patients were further categorized based on the presence of joint pain as Pre-Menopausal Arthralgia (PrMA), Peri-Menopausal Arthralgia (PeMA) and Post-Menopausal Arthralgia (PoMA) [15]. The definition of menopause included cessation of menses for at least 1 year, follicle stimulating hormone (FSH) levels of ≥ 40 mIU/ml and <20 pg/ml of estradiol [16]. Each patient's age and baseline joint pain score (visual analog scale: VAS 100 mm in length anchored by 2 verbal descriptors [17]) was documented on the first visit and at subsequent visits. Any reduction of VAS was noted. FSH and estrogen levels were determined on the first visit, then after two months and semi-annually thereafter.

Hormone replacement therapy

Premenopausal women were treated with ultra-low dose norethisterone (1 mg/ethinylestradiol 0.035 mg), PeMA women with α -tocopherol N 600 mg and postmenopausal women with HRT for at least two months, respectively. For women less than 45 or over 60 and those with contraindications to HRT, HRT was not used. The same therapy was continued in patients who experienced a good response. Estrogen monotherapy (i.e., estrogen replacement therapy (ERT)) was administered to PoMA women with a previous hysterectomy. In females without a hysterectomy, monthly treatment with a 17β -Estradiol patch (0.72 mg)/2 days for 26 days and dydrogesterone (duphaston) 10 mg for

10 days, followed by 5 days off the treatment was administered. In PoMA patients ≥ 5 years, estradiol 1mg/levonorgestrel 0.04 mg) 1 tablet/day was prescribed as EPRT continuous. In patients ≥ 60 years of age, estriol (1 mg) 2 tablets/day was prescribed and dydrogesterone 10 mg $\times$ 10 days were taken every 6 months.

ANA and anti-DFS70 autoantibody determination

Serum ANA was detected by IIF on HEp-2 cells (Medical and Biological Laboratories Co. Ltd. (MBL), Nagoya, Aichi, Japan). A detectable cellular staining pattern at a serum dilution of $\geq 1:40$ was regarded as positive. Anti-DFS70 Aab was measured by enzyme-linked immunosorbent assay (ELISA, MBL) and the cutoff of 15 units was established according to the manufacturer's instructions.

Statistical methods

Statistical analysis included the frequency of anti-DFS70 in all SARD and non-SARD PrMA vs. PeMA vs. PoMA women; the frequency of anti-DFS70 in SARD vs. non-SARD PeMA women; the correlation between ANA IIF and anti-DFS70 Aab. The Mann-Whitney U test and χ^2 -tests were used for clinical significance.

RESULTS

Baseline demographics and serological profile

The menopausal status and diagnostic distribution of the 282 Japanese females enrolled in this study are shown in **Figure 1**. Their age range was 29-91 years (median=58). 122 had UA, 116 had a SARD, 28 had 'other' diagnoses and 16 were

healthy individuals. Out of the patients with UA, 17 were pre-menopausal, 21 were peri-menopausal, 84 were post-menopausal.

A positive ANA was found in 84.1% (237/282) patients. Anti-DFS70 Aab was found in 17.4% (49/282) of women (**Figure 1a**) and was detected in all age groups (**Figure 2**).

When females younger than 30 and older than 80 years were excluded because of small numbers, more than 50% of positive anti-DFS70 Aab subjects were in the in 46 to 59 year old cohort. Frequencies of anti-DFS70 Aab in 45-49, 50-54 and 55-59 age groups were 18.5% (5/27), 25.5% (12/47) and 14.2% (7/49), respectively (Chi square test $p=0.52$, not statistically significant).

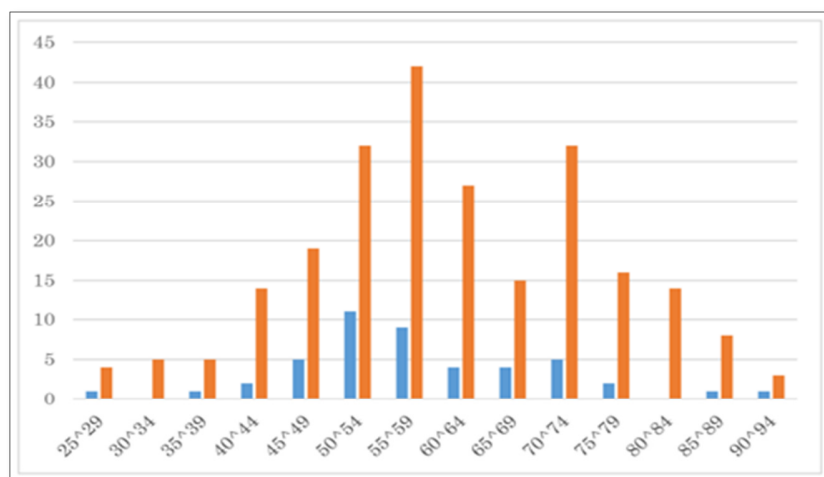


Figure 2. Anti-DFS70 Aab in Japanese females at different ages (5 year intervals)*.

* positive anti-DFS70 positive/number of patients in each 5 year interval: 1/5 0/5 1/6 2/16 5/24 11/43 9/51 4/31 4/19 5/37 2/18 0/14 1/9 1/4

Frequency and anti-DFS70 Aab titers in pre-menopausal, peri-menopausal and post-menopausal women with UA

The frequency of anti-DFS70 Aab in 122 UA women was 17.6% (3/17) with PrMS, in 28.5% (6/21) with PeM and in

26.1% (22/84) with PoM (**Table 1**). The mean anti-DFS70 ELISA titers (absorbance index) of 8, 16.9 and 15.7 were observed in PrMS, PeMS and PoMS, respectively.

Table 1. Frequency and anti-DFS-70A antibody titers in pre-menopausal, peri-menopausal and post-menopausal women.

	PrMA	PeMA	PoMA	RA	SjS	SLE	SSc	MCTD	HI
Patient N	17	21	84	65	31	10	6	4	15
Positive N	3	6	22	7	3	0	0	0	2
Positive %	17.6	28.5	26.1	10.8	9.7	0	0	0	13.3
Mean Index	8	16.9	15.7	8.1	4.9	1.8	1.7	2.6	16.8
Mean age	33.6	48.4	56.9	66.4	66.1	60.8	74.3	48	51.9
E2 (pg/mL)	166.2 (1)*	88.1 (10)*	32.3 (27)*	nd	nd	nd	nd	nd	nd
FSH (mIU/mL)	5.3 (1)*	42.3 (10)*	47.2 (27)*	nd	nd	nd	nd	nd	nd

Abbreviations: E2: Estradiol; FSH: Follicle Stimulating Hormone; MCTD: Mixed Connective Tissue Disease; nd: not done; PeMA: Peri-Menopausal Arthralgia; PoMA: Post-Menopausal Arthralgia; PrMA: Pre-Menopausal Arthralgia; RA: Rheumatoid Arthritis; SjS: Primary Sjögren's Syndrome; SLE: Systemic Lupus Erythematosus; SSc: Systemic Sclerosis; UA: Undifferentiated Arthralgia/Arthritis

* = number tested

Note: Twenty-nine women including 7 with other miscellaneous diseases (**Figure 1**) were excluded. Serum E2 and FSH levels performed when anti-DFS70 Aab were tested

The frequency of anti-DFS70 Aab of 26.7% (28/105) in all PeM and PoM UA females was significantly higher than that of 10.8% in women with RA or 9.7% in women with primary SjS ($p=0.013$ by U test). Further, the mean ELISA index of 16.1 in PeMA and PoMA females was significantly higher than 8.1 in RA and 4.9 in primary SjS, respectively (**Table 1**). By comparison, 2 of 15 (13.3%) HI had anti-DFS70 Aab and the observed frequencies of anti-DFS70 of 28.5% in UA PeMA and 26.1% in UA PoMA were higher than HI (not statistically significant) than in the other menstrual stages (PrMSA) studied here.

Anti-DFS70 Aab in pre-, peri- vs. post-menopausal women progressing to SARDs vs. non-SARD

The frequency of anti-DFS70 Aab of 41.2% (7/17) in PoMA associated with SARD was significantly higher than that the 14.7% (15/67) in PoMA without SARD ($p<0.05$) (**Table 2**). In addition, although the number of subjects is small, the frequency of anti-DFS70 Aab of 66.7% (2/3) in PeMA associated with ongoing to SARD is higher than that with 26.6% (4/18) in PeMA without SARD. The frequency of anti-DFS70 Aab of 37.5% (3/8) in PrMA with SARD is higher than that of 0% in 9 women without SARD (Correlation coefficient: $y=0.1326x + 190.53$; $r=0.009$).

Table 2. Frequency and mean ELISA OD index of anti-DFS70 Aab in pre-, peri- and post-menopausal women with and without association of SARD treated with HRT.

	Total	Non-SARD				N	SARD		
		N	Positive	%	Mean Index		Positive	%	Mean Index
PoMA	84	67	15	14.7	22.4	17*	7	41.2	19.7
PeMA	21	18	4	26.6	16.2	3**	2	66.7	20.6
PrMA	17	9	0	0	1.5	8***	3	37.5	15.4

Abbreviations: SARD Systemic Autoimmune Rheumatic Disease; Poma: Post-Menopausal Arthralgia; Pema, Peri-Menopausal Arthralgia; Prma: Pre-Menopausal Arthralgia
SARD include:

* RA (2), possible RA (1), SjS (3), probable SjS (4), SLE (1)

**RA (1), SjS (1), possible SjS (1)

***RA (2), possible RA (1), SjS (2), SLE (1), MCTD (2)

Non-SARD include: FM (3), autoimmune hepatitis (1), Hashimoto thyroiditis (1), severe OA and possible RA (1)

Relationship between the ANA IIF titers of and anti-DFS70 ELISA among all 282 enrolled Japanese women

Anti-DFS70 Aab were found in 49/282 (17.3%) of all patients and in 49/237 (20.7%) of ANA positive patients (**Figure 3a**). There was a tendency for high titer ANA to be associated with low titer anti-DFS70 Aab by ELISA, and ANA low titer with high anti-DFS70 by ELISA (**Figures 3a and 3b**). Statistical analysis between ANA IIF titres and DFS70 antibodies revealed a correlation coefficient of $r=0.357$. In assessing ANA-related diseases, higher ANA IIF titers ($\geq 1:320$) were associated with lower anti-DFS70 Aab

titers in 3 of 10 (30%) SLE and 15 of 31 (48.4%) SjS patients.

The majority of anti-DFS70 Aab positive sera had a positive ANA at dilutions of 1:40-1:160. For example, 40/49 (81.6%) ANA positive with anti-DFS70 Aab sera showed anti-DFS (homogeneous/speckled) IIF staining patterns at the IIF screening dilution of $\geq 1:40$ (**Figure 3b**). The remainder of the IIF patterns included nuclear envelope ($n=1$), speckled ($n=2$), discrete speckled ($n=2$), PCNA-like ($n=1$) and negative ($n=3$).

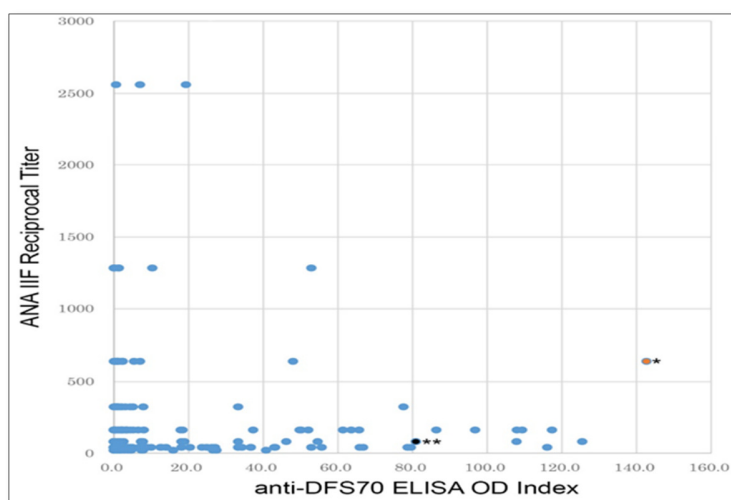


Figure 3a. Relationship between the ANA IIF titers of and anti-DFS70 ELISA among all 282 enrolled Japanese women. ANA is negative at <40 and anti-DFS70 Aab is negative at ≤ 15 OD index. Women designated * and ** are the results of the anti-DFS assays of two case reports. There is a tendency of high titer ANA to be associated with low titer anti-DFS70 Aab by ELISA and ANA low titer and high anti-DFS70 by ELISA ($y=0.1326x + 190.53-r=0.009$)

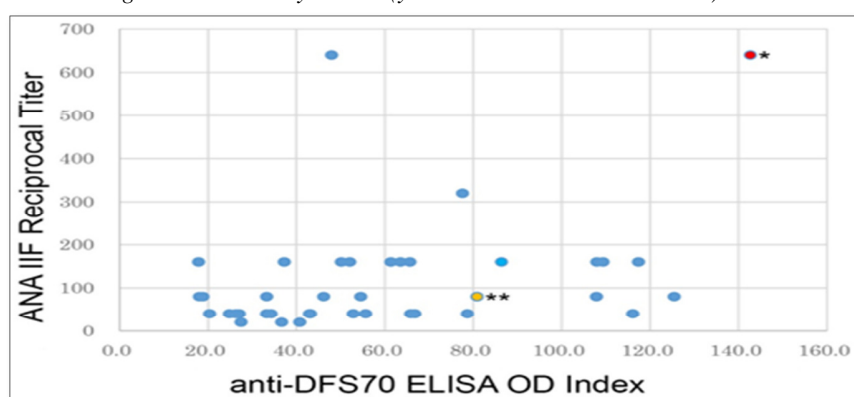


Figure 3b. Relationship between ANA IIF titers and anti-DFS 70 Aab as detected in the 40 sera with a staining pattern consistent with DFS and an anti-DFS70 antibody positive by ELISA (OD Index > 15) (some plots overlap). Single plots denoted by * and ** represent the case reports of Patients 1 and 2, respectively

DISCUSSION

We began with a presentation of two postmenopausal patients presenting with arthralgias in the context of a positive ANA by IIF and monospecific anti-DFS70 antibody. Despite a positive ANA, the rheumatologists did not find evidence for a SARD and their arthralgia was thought to be secondary to their post-menopausal state. Their symptoms were treated successfully with cyclic HRT for 2 months. Menopausal arthralgias not well understood, however, they are very common complaints in Japan, yet only 2~3% of these patients received HRT compared to 20 to 50% of menopausal women receiving HRT in USA, Europe and Australia [18]. HRT treatment in some jurisdictions in Japan is administered by general practitioners, gynecologists and internists, but uncommonly rheumatologists.

Our patients with menopausal joint pains achieved less than 5 points in the ACR/EULAR classification criteria for early RA and hence were classified as undifferentiated arthritis (UA) [11]. As previously reported in a Dutch study, over a one year follow-up, 40% of UA developed RA and 20% UA progressed to defined diseases such as Sjögren's syndrome and 40% UA spontaneously resolved [14]. One of the prevailing concerns of rheumatologists is the use of an ANA in patients with 'aches and pains' (i.e., as in our patient 1) to rule out a diagnosis of SARD [19]. However, it is precisely in this setting that the ANA can be helpful because if it is found to be monospecific anti-DFS70 it effectively rules out the diagnosis of a SARD [20,21] and can help allay patient anxiety and physician uncertainty [4,5]. Since HRT is not so effective in treating overt RA [22], it may be helpful in discriminating estrogen deficiency or depletion arthralgia (menopausal arthralgia) from early stage of RA. For

example, in our experience HRT treatment of RA results in less than 30% reduction of the visual analog joint pain scale (unpublished).

In general, reports to date indicate that the frequency of anti-DFS70 is higher in younger females [23] and in apparently healthy individuals and individuals with SARD than it is in individuals with a SARD [4,8,24]. In our study, anti-DFS70 Aab was found in 16.3% of apparently healthy Japanese women, which is comparable to the frequency reported by Watanabe et al. [9], who, using an immunoblotting and ELISA, reported a frequency of 10.7% in a study of healthy Japanese hospital workers, 75.9% of which were females with a mean age of 31 years. In our study there was no significant difference in the frequency of anti-DFS70 Ab in females with UA at different stages of menopause as compared to HI and the frequency of anti-DFS70 Aab in females with UA associated with menopausal complaints (26.7%) was significantly higher than those with possible/probable or defined SARD (9.7% primary SjS; 10.8% RA). These observations are inconsistent with the study of Watanabe who reported anti-DFS Aab in only 1.5% of individuals with a SARD [25]. Since anti-DFS70 is found in healthy individuals as well as SARD, further study has shown that a key differentiating feature is that in healthy individuals the anti-DFS70 response tends to be restricted to that antigen whereas in SARD, the Aab are found in association with other SARD-related Aab [26]. In our two patients reported here no other SARD-related Aab were found.

The hypothesis that the estrogen hormonal status may be an explanation for anti-DFS70 B-cell responses was not proven by our study where the focus was on pre-, peri- or post-menopausal women who lacked a SARD. The anti-DFS70 frequency of ~27% in PeMA and PoMA patients in our study is among the highest published in adult or pediatric cohorts to date. Previous studies reported that the frequency of anti-DFS70 may be higher in younger females [9,23], especially in pre-pubertal children [10]. Although there was a trend to a lower frequency of anti-DFS in PrMA, our PrMA numbers were small because of the focus on menopausal women rather than PrMA and the youngest PrMA patient was 26 years of age.

A limitation of our study is that we did not determine if there was a correlation between anti-DFS70 Aab levels and estrogen/FSH levels. In addition, since our study did not focus on young or pre-menopausal Japanese females without arthralgia or SARD, further study on this population is required. In addition, earlier studies have reported anti-DFS Aab in in 3~30% of atopic dermatitis or alopecia [27] and in 10% of interstitial cystitis [6]. Moreover, it was found in ~30% of children with a positive ANA and chronic fatigue syndrome [28] and in patients with cataracts [7]. However, in our study, we did not systematically evaluate patients for these conditions. Although it has been reported that anti-

DFS70 responses remain relatively stable over time [29,30], formal, well designed longitudinal studies are still required.

In our two case reports, we reported the ANA IIF description that was provided in the laboratory diagnostic report (homogenous and speckled). The ICAP nomenclature for DFS immunofluorescence pattern on HEp-2 cells (AC-2) has only recently been described on the website (<https://anapatterns.org/index.php>). Nevertheless, many laboratories do not report the DFS IIF pattern as described and one recent study showed that there continues to be significant inter-laboratory disparity in recognizing and reporting the DFS IIF pattern [31]. Indeed, if one compares the ICAP AC-1 and AC-2 patterns and descriptors they are very similar in that both stain the interphase nuclei and the metaphase chromatin. We confirmed the DFS (AC-2) pattern for both Patient 1 and Patient 2.

In our study, anti-DFS70 Aab was most commonly found in sera with ANA IIF titers of less than 1:160. It is well-known that the titers of anti-DFS70 as detected by IIF can be very high (i.e., >1/2560) [32]. ANA titers of 1:320 or greater were also associated with anti-DFS70 Aab in our study. However, only 6 of all 46 anti-DFS70 positive sera were positive for anti-DFS70 Aab (ELISA index >15) at an ANA of 1:320 or greater. These data suggest that anti-DFS70 Aab as detected by our ELISA that utilized a human recombinant protein is not highly correlated with conventional ANA IIF patterns, a finding that is consistent with previous reports showing that not all patients with anti-DFS70 have the complimentary DFS IIF staining pattern on HEp-2 cells [26,33].

Since anti-DFS70 Aab may be associated with the HLADRB1 genetic background [34] that along with infections, stressor proteins and estradiol fluctuation or depletion might further enhance the autoimmune response [35]. Therefore, in addition to defining HLADRB1 status, future studies might survey patients for infections (i.e., microbiome) or stressor proteins [36] as well as hormonal status as possible contributors to the anti-DFS70 response. In addition, considering the possible role of hormones like estrogen on autoimmune responses, follow-up anti-DFS70 and ANA testing after HRT would be a potentially informative study. In this context, it is possible that menopausal related hormonal changes may be associated with viral reactivation and transient arthralgia and an anti-DFS70 response. Ongoing studies are hoping to determine if a viral (or bacterial) exposure triggers and perpetuates the anti-DFS70 response. Based on demographic data alluded to above; potential triggers might include Epstein Bar virus or the human papilloma virus. However, the human immunodeficiency virus may be more relevant because the N-terminus of DFS70 plays a role in the recruitment of the cognate DFS70 protein to chromatin and the C-terminus contains a highly conserved domain termed the human immunodeficiency virus integrase (HIV-IN) binding domain

(IBD) that promotes the integration of HIV DNA into transcriptionally active chromatin [37]. Of note, the IBD also represents the predominant DFS70 epitope that binds human anti-DFS70 antibodies [37].

In summary, an important contribution of our study and supporting case reports is that in the management of menopausal females where joint pains can be a significant clinical component, an assessment of anti-DFS70 is an important consideration. In these patients, the finding of anti-DFS70 is an important step to ruling out a SARD and these patients can benefit from HRT. Further, our studies also support the evidence that the reliable detection of anti-DFS70 requires testing for anti-DFS70 Aab using ELISA or other analyte-specific assays [4,38]. If positive, unless the clinical picture changes, they should not continue visiting rheumatology clinics for evaluation SARD as part of a cost-effective approach to diagnostics [35].

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CONFLICTS OF INTEREST

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The other authors have no conflict of interest.

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