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1	Article Title	<b>Chronic high-dose glucocorticoid therapy triggers the development of chronic organ damage and worsens disease outcome in systemic lupus erythematosus</b>	
2	Article Sub- Title		
3	Article Copyright - Year	<b>International League of Associations for Rheumatology (ILAR) 2016 (This will be the copyright line in the final PDF)</b>	
4	Journal Name	Clinical Rheumatology	
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44		e-mail	
45		Received	7 October 2016
46	Schedule	Revised	10 November 2016
47		Accepted	20 November 2016
48	Abstract	Long-term survival of patients with systemic lupus erythematosus (SLE) improved worldwide; thus, prevention of cumulative organ damage became a major goal in disease management. The aim of our study was to investigate the chronic organ damages and their influence on disease outcome in SLE. We evaluated clinical conditions, laboratory findings and medications of 357 consecutive SLE patients and assessed their impact on Systemic Lupus Collaborating Clinics (SLICC)/American College of Rheumatology (ACR) Damage Index (SDI) and disease outcome. We detected one or more SDI scores in 77.87% of patients. Patients with disease duration of more than 10 years and subjects diagnosed at age above 40 had significantly higher SDI values. The most frequent damages were valvulopathies, cognitive dysfunction, angina pectoris and venous thrombosis. Higher cumulative glucocorticoid dose increased SDI, while chloroquin treatment was favourable for patients. Male gender, elevated SDI scores and higher cumulative doses of glucocorticoids increased mortality risk. Our data confirmed that disease duration, age at diagnosis and chronic high-dose glucocorticoid therapy have significant effects on the development of chronic organ damage. Higher SDI score is characterized with worse survival ratios. The most common chronic	

organ damages affected the cardiovascular or neuropsychiatric system. As long-term survival in SLE improves, it becomes increasingly important to identify the determinants of chronic organ damage. Most of the chronic organ damage occurs in the cardiovascular and the neuropsychiatric systems; thus, regular follow-up, screening and adequate therapy are essential for the best clinical outcome.

- 
- 49 Keywords                      Chronic organ damage - Disease outcome - SLICC/ACR Damage  
separated by ' - '              Index - Systemic lupus erythematosus
- 
- 50 Foot note  
information

# Chronic high-dose glucocorticoid therapy triggers the development of chronic organ damage and worsens disease outcome in systemic lupus erythematosus

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9 Received: 7 October 2016 / Revised: 10 November 2016 / Accepted: 20 November 2016  
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16 come in SLE. We evaluated clinical conditions, laboratory  
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32 organ damage. Higher SDI score is characterized with worse  
33 survival ratios. The most common chronic organ damages  
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35 long-term survival in SLE improves, it becomes increasingly  
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38 vascular and the neuropsychiatric systems; thus, regular

follow-up, screening and adequate therapy are essential for  
the best clinical outcome. 39 40

**Keywords** Chronic organ damage · Disease outcome · 41  
SLICC/ACR Damage Index · Systemic lupus erythematosus 42

## Introduction 43

44 Systemic lupus erythematosus (SLE) is a chronic autoimmune 44  
45 disease that can affect almost any organs and tissues of the 45  
46 body, leading to a wide spectrum of clinical manifestations. 46  
47 For a long time, lupus was considered to be a disease with a 47  
48 poor prognosis, but in recent years, the long-term survival in 48  
49 SLE has improved significantly. While during the 1960s, the 5- 49  
50 year survival rate was 60%, by the 2000s, it has increased up to 50  
51 90% in most countries and centres [1, 2], although ethnic and 51  
52 geographic variations remained significant [3, 4]. However, the 52  
53 increased longevity of patients with SLE leads to the accumu- 53  
54 lation of chronic organ damage over time in patients, which 54  
55 became one of the most important factors that contribute to 55  
56 mortality in SLE [5]. Disease activity and certain comorbidities 56  
57 are the main factors; however, several other factors are known 57  
58 to influence the development of chronic organ damage. 58  
59 Importantly, immunomodulatory treatments can be also associ- 59  
60 ated with adverse events, organ damages and mortality. La 60  
61 Gonzales et al. identified menopause as well as gender, age 61  
62 and ethnicity as further significant influencing factors; more- 62  
63 over, they reported that certain psychosocial factors can also 63  
64 promote chronic damage [6]. Therefore, it is important to ex- 64  
65 amine and understand the factors and mechanisms that influ- 65  
66 ence disease prognosis and patients' quality of life. 66

67 The Systemic Lupus Collaborating Clinics (SLICC) and the 67  
68 American College of Rheumatology (ACR) proposed the inter- 68  
69 nationally validated damage scoring system, namely, SLICC/ 69

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70 ACR Damage Index (SDI) for the evaluation of chronic organ  
 71 damage. SDI can be used to measure the degree of damage and  
 72 to check its change over time [7]. Previous studies revealed  
 73 significant associations between damage; disease activity; and  
 74 certain demographic, clinical and laboratory features [8, 9].

75 Due to lack of data from East-Central Europe, the aims of  
 76 our work were to survey SDI values in a large cohort of  
 77 Hungarian SLE patients, to compare our results with interna-  
 78 tional data and to identify additional influencing factors.

79 **Material and methods**

80 **Patients**

81 In our present cross-sectional study, we evaluated 357  
 82 Hungarian patients with SLE who were diagnosed between  
 83 1 January 1971 and 31 December 2012 and also treated at the  
 84 Division of Clinical Immunology in the Medical Center of  
 85 University of Debrecen. All patients were followed up on a  
 86 routine basis, and their records contained detailed information  
 87 on symptoms, clinical conditions, laboratory and other find-  
 88 ings of each visit. The diagnosis of SLE was established based  
 89 on the ARA preliminary classification criteria or ACR classi-  
 90 fication criteria revised in 1982 or in 1997, according to the  
 91 date of first visit [10–12]. Patients diagnosed with SLE before  
 92 1997 were revised according to the revised 1997 ACR criteria  
 93 for SLE classification. Sapporo and Sydney criteria were used  
 94 to establish the diagnosis of anti-phospholipid syndrome [13,  
 95 14]. All experiments carried out in the study were in compli-  
 96 ance with the Declaration of Helsinki.

97 **Clinical evaluations**

98 All patients were followed up on a routine basis, and their  
 99 records contained detailed information on symptoms, clinical  
 100 conditions and laboratory and other findings of each visit. The  
 101 following demographic and clinical data were analyzed: gen-  
 102 der, age, age at diagnosis, duration of disease, clinical symp-  
 103 toms and organ manifestations of SLE, comorbidities, labora-  
 104 tory parameters, immunoserological abnormalities and thera-  
 Q1 105 py used during the disease course. Disease activity was mea-  
 106 sured using Systemic Lupus Erythematosus Disease Activity  
 107 Index (SLEDAI) [15, 16]; flare was defined as an increase in  
 108 SLEDAI score with at least 3 points. The assessment of chron-  
 109 ic organ damage was performed using SDI [7].

110 **Laboratory measurements**

111 Immunoserological tests were performed at the Regional  
 112 Immunology Laboratory of the Division of Clinical  
 113 Immunology and included the measurement of anti-  
 114 nuclear antibody (ANA), rheumatoid factor (RF),

antibodies against extractable nuclear antigen (ENA), anti- 115  
 anti-dsDNA, anti-Sm, anti-RNP, anti-SS-A, anti-SS-B, anti- 116  
 phospholipid antibodies, serum immunoglobulins, 117  
 haemolysis test and complement levels. Hep-2 cell-based 118  
 indirect immunofluorescence assay was performed as a 119  
 screening test for anti-ENA antibodies, and further iden- 120  
 tification was carried out by enzyme-linked immunosor- 121  
 bent assay (ELISA) with AUTOSTAT II kits (Hycor 122  
 Biomedical, Indianapolis, IN, USA), according to the 123  
 manufacturer’s instructions. Immunoglobulin levels and 124  
 complement activity were determined with turbidimetry 125  
 and nephelometry techniques and haemolysis test in sheep 126  
 red blood cell suspension, respectively. General laboratory 127  
 parameters (blood count, kidney and liver function, 128  
 haemostasis parameters, lupus anti-coagulant, urinalysis) 129  
 were assessed at the Clinical Biochemistry and Molecular 130  
 Pathology Institute of University of Debrecen. 131

**Therapy** 132

We registered the use of medications, including glucocorti- 133  
 coids, immunosuppressive agents, hydroxychloroquine and 134  
 biologics. Additionally, we also calculated the cumulative 135  
 dosage of glucocorticoids and analyzed the relationship be- 136  
 tween SDI and the different treatment modalities. 137

**Statistical analyses** 138

The IBM SPSS ver. 22.0 (SPSS Inc., Chicago, IL, USA) 139  
 was used for statistical analysis. In cases of continuous 140  
 variable, we determined mean and standard deviation 141  
 (SD) values and used independent samples *t* test or 142  
 Mann-Whitney test for statistical evaluation. When the 143  
 strength of the linear relationship between two variables 144  
 was evaluated, Pearson’s correlation coefficient was used, 145  
 while in cases of non-normal distribution, Spearman’s cor- 146  
 relation coefficient was applied. Chi-square test and 147  
 Fisher’s exact test were used to discriminate between pa- 148  
 tient groups. Data on disease outcome are given in mean 149  
 values with 95% confidence intervals (CIs). We used the 150  
 Cox regression model to predict chronic organ develop- 151  
 ment in the disease. Survival time and rate were assessed 152  
 using the Kaplan-Meier estimator. Chi-square test and 153  
 Fisher’s exact test were used to discriminate between pa- 154  
 tient groups, and we used the Cox regression model to 155  
 predict poor outcome of the disease. Differences were con- 156  
 sidered statistically significant at  $p < 0.05$ . 157

**Results** 158

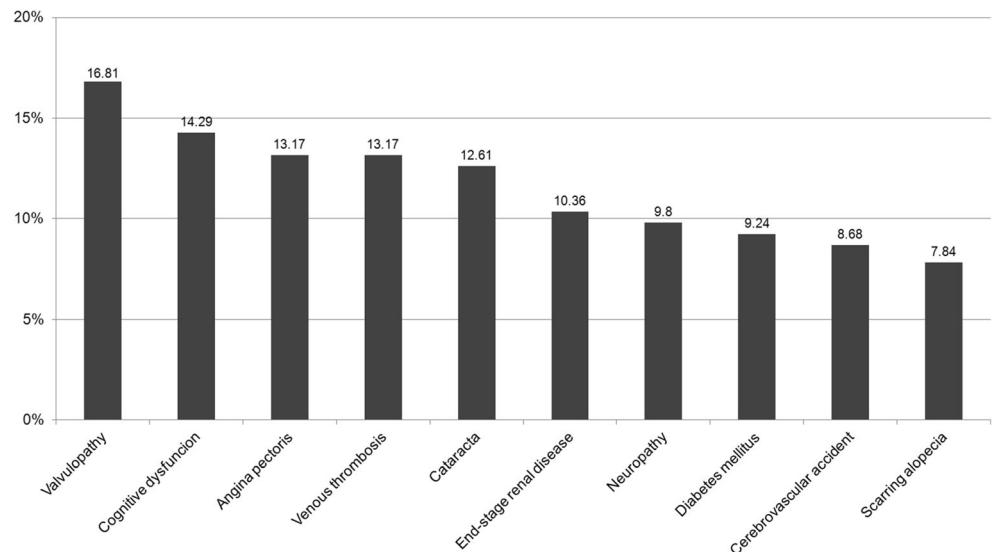
Table 1 summarizes the demographic data and clinical and lab- 159  
 oratory features of the 357 SLE patients. The mean follow-up 160

t1.1 **Table 1** The main demographic, clinical and serological features of SLE patients (*n* = 357)

t1.2	Demographic features	
t1.3	Male/female	33/324
t1.4	Age (years) mean ± SD (range)	51.57 ± 13.48 (21–86)
t1.5	Age at disease onset (years) mean ± SD (range)	32.11 ± 11.49 (7–67)
t1.6	Disease duration (years) mean ± SD (range)	19.14 ± 9.15 (1–44)
t1.7	Clinical damages, <i>N</i> (%)	
t1.8	Cardiovascular damage	108 (30.25)
t1.9	Neuropsychiatric damage	91 (25.49)
t1.10	Musculoskeletal damage	65 (18.21)
t1.11	Peripheral vascular damage	57 (15.97)
t1.12	Ocular damage	56 (15.68)
t1.13	Renal damage	56 (15.68)
t1.14	Skin damage	50 (14.01)
t1.15	Pulmonary damage	35 (9.8)
t1.16	Gastrointestinal damage	3 (0.84)
t1.17	Serological abnormalities, <i>N</i> (%) last time of the follow-up	
t1.18	ANA	355 (99.44)
t1.19	Anti-dsDNA	195 (54.62)
t1.20	Anti-Sm	86 (24.09)
t1.21	Anti-SSA	99 (27.73)
t1.22	Anti-SSB	59 (16.53)
t1.23	Anti-cardiolipin IgG/IgM	86 (24.09)
t1.24	Anti-beta2 GPI IgG/IgM	75 (21.01)
t1.25	Lupus anti-coagulant	24 (6.72)
t1.26	Low C3/C4	153 (42.86)
t1.27	Medications, <i>N</i> (%)	
t1.28	Glucocorticoids	310 (86.83)
t1.29	Cumulative dosage of glucocorticoids (g) mean ± SD	32.878 ± 25.506
t1.30	Chloroquine	158 (44.26)
t1.31	Azathioprine	171 (47.9)
t1.32	Cyclophosphamide	103 (28.85)
t1.33	Methotrexate	40 (11.2)
t1.34	Biologics	36 (10.08)
t1.35	Cyclosporine A	21 (5.88)
t1.36	Leflunomide	16 (4.48)
t1.37	Mycophenolate mofetil	12 (3.36)

161 period was 19.14 ± 9.15 years with a range 1 to 44 years. The  
 162 mean age of patients at the time of their last follow-up visits was  
 163 51.57 ± 13.48 years with a range 21 to 86 years, while their  
 164 mean age at disease onset was 32.11 ± 11.49 years (range 7–  
 165 67 years). There were 33 male (9.24%) and 324 female  
 166 (90.76%) patients; male to female ratio was 9.8:1.

**Fig. 1** Percentages of the ten most frequent specific chronic organ damages in SLE patients (*n* = 357)



**Chronic organ damage**

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Based on our observations, men had higher mean SDI value (SDI: 2.03 ± 1.55) compared to women (SDI 1.88 ± 1.73), but the difference was not significant.

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Out of 357 patients, 278 patients (77.87%) were found to have developed at least one chronic organ damage. Damage scores 1 and 2 were the most frequent [*N* = 104 (29.13%) and *N* = 62 (17.37%), respectively], followed by scores 3 and 4 [*N* = 56 (15.69%) and *N* = 25 (7%), respectively] and scores 5 and 6–8 [*N* = 15 (4.2%) and *N* = 16 (4.48%), respectively]. The cardiovascular organ system was the mostly affected in patients during the disease course (*N* = 108, 30.25%). Ninety-one patients (25.49%) were found to have developed neuropsychiatric, 65 patients (18.21%) musculoskeletal and 57 patients (15.97%) peripheral vascular, and both ocular and renal damage affected 56 patients (15.68%). Fifty patients (14.01%) were found to have dermatological, 35 patients (9.8%) pulmonary and 3 patients (0.84%) gastrointestinal organ system damage (Table 1). The ten most frequent types of chronic organ damage are listed in Fig. 1.

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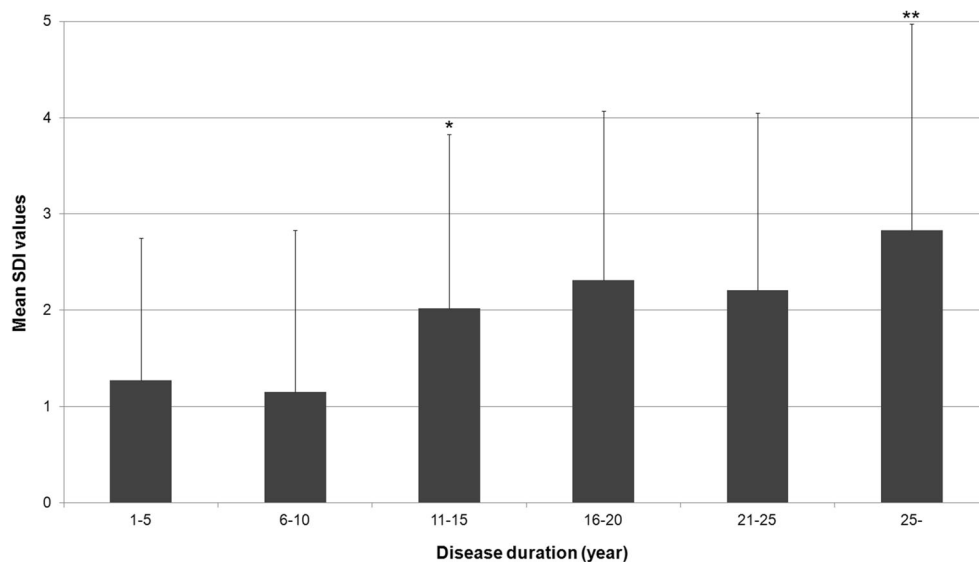
Based on our results, the number of chronic damages was significantly higher in patients with disease duration of more than 10 years (mean SDI value of patients with disease duration of 6–10 years, 1.15 ± 1.68 vs. mean SDI value determined in patients with disease duration of 11–15 years, 2.02 ± 1.81, respectively, *p* = 0.014). Patients with a disease duration of more than 25 years had even higher SDI values (mean SDI value of patients with disease duration of 21–25 years, 2.21 ± 1.84 vs. mean SDI value determined in patients with disease duration of more than 25 years, 2.83 ± 2.14, respectively, *p* = 0.018) (Fig. 2).

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We examined the relationship between the SDI value and disease activity, as well. Of patients without chronic damage, 25.32% developed a disease flare during the last 10 years of the study. Of patients with a score of 1–3, 28.63% showed

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**Fig. 2** Association between the disease duration and SDI. Patients with disease duration of more than 10 years had higher SDI values ( $*p = 0.014$ ). Patients with disease duration of more than 25 years had even higher SDI values ( $**p = 0.018$ )



201 active disease. Of patients with an SDI value of at least 4,  
 202 32.29% demonstrated disease flare. The increase in SDI  
 203 values was mirrored by an increase in the number of patients  
 204 with disease flare, but the difference was not significant.

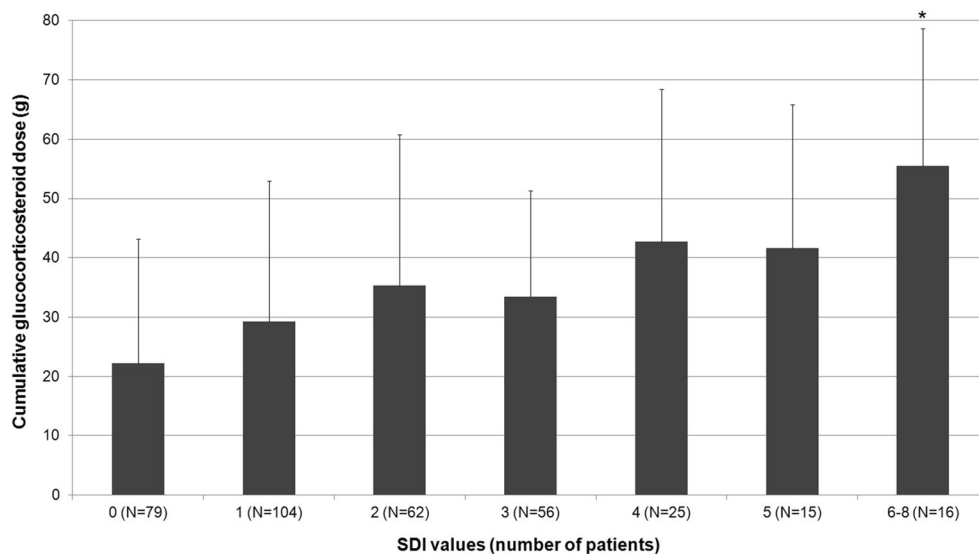
205 The patients' mean age at diagnosis had an influence on the  
 206 SDI value. The SDI value of SLE patients who were diag-  
 207 nosed above the age of 40 years ( $N = 102$ ) was significantly  
 208 higher than the mean SDI value of patients diagnosed under  
 209 40 years ( $N = 255$ ) ( $2.28 \pm 1.92$  vs.  $1.74 \pm 1.6$ , respectively,  
 210  $p = 0.007$ ).

211 We also investigated the relationship between SDI and the  
 212 different treatment modalities. Regarding long-term glucocor-  
 213 ticoid therapy, patients with a higher SDI score (6–8) had a  
 214 significantly higher ( $p < 0.001$ ) cumulative glucocorticoid  
 215 dose than patients with lower SDI scores (1–2). Patients who  
 216 received higher-dose glucocorticoid therapy had higher mean  
 217 SDI scores (Fig. 3). Furthermore, significantly higher average

218 cumulative glucocorticoid dose was administered to SLE pa-  
 219 tients with cataracts ( $p < 0.001$ ) or osteoporosis ( $p = 0.041$ ).  
 220 Cumulative doses were also higher in patients with cerebro-  
 221 vascular events, lower extremity claudication, myopathy and  
 222 avascular necrosis of the femoral head, but the difference was  
 223 not statistically significant. We also revealed a strong positive  
 224 correlation between SDI values and cumulative glucocorticoid  
 225 doses in the whole cohort of SLE patients ( $R = 0.307$ , respec-  
 226 tively,  $p < 0.001$ ). Moreover, adjusted odds ratios (ORs) by  
 227 multiple logistic regression analysis showed that cumulative  
 228 doses were significantly and independently related to SDI  
 229 (OR 0.05, respectively,  $p = 0.027$ ).

230 Interestingly, the mean SDI value of patients treated with  
 231 chloroquine ( $N = 158$ ) was significantly lower than that of  
 232 lupus patients not receiving chloroquine ( $1.64 \pm 4.54$  vs.  
 233  $2.1 \pm 1.82$ , respectively,  $p = 0.024$ ). In the cases of cyclophos-  
 234 phamide, azathioprine, methotrexate, cyclosporine A and

**Fig. 3** The effect of long-term glucocorticoid therapy on SDI values. Patients with the highest SDI values (6–8) had a significantly higher average cumulative glucocorticoid dose compared to patients with lower SDI values (0–5) ( $*p < 0.001$ )





235 other investigated therapies, there was no significant differ- 267  
 236 ence between the mean SDI values of treated and non- 268  
 237 treated patients. We did not find any associations between 269  
 238 serological parameters and SDI values. 270

239 **Disease outcome**

240 During the whole follow-up period, 42 (32 women and 10 men) 275  
 241 of our patients died. Mortality of the whole patient population 276  
 242 was 11.76%; of note, mortality values differed significantly 277  
 243 between male and female patients (30.3 vs. 9.88%, respectivel- 278  
 244 y,  $p = 0.002$ ). As to the distribution by age groups, we lost 20 279  
 245 (17 female and 3 male), 18 (13 female and 5 male) and 4 (2 280  
 246 female and 2 male) patients, from the >60 years, the 40– 281  
 247 59 years and the <40 age groups, respectively. When evaluating 282  
 248 the causes of death, infections ( $N = 15$ ) and cardiovascular 283  
 249 events, such as myocardial infarction ( $N = 11$ ) and stroke 284  
 250 ( $N = 3$ ), were the leading causes, being followed by heart failure 285  
 251 ( $N = 3$ ) and tumours including lung ( $N = 3$ ), breast ( $N = 2$ ), liver 286  
 252 ( $N = 1$ ) and brain cancers ( $N = 1$ ), as well as malignant mela- 287  
 253 noma ( $N = 1$ ) and non-Hodgkin's lymphoma ( $N = 2$ ). 288

254 The overall 5-year survival rate was 99%, the 10-year sur- 289  
 255 vival rate was 98%, and the 15-year survival rate was 95%. 290  
 256 The mean survival was 37.21 years [95% confidence interval 291  
 257 (CI), 35.33–39.1]. Male patients and patients with 5 or more 292  
 258 SDI score could be characterized with significantly worse sur- 293  
 259 vival ratios. The mean survival of male patients was signifi- 294  
 260 cantly worse, compared to the values of female patients 295  
 261 [28.78 years (95% CI, 24.82–32.74) vs. 38.19 years (36.24– 296  
 262 40.15), respectively,  $p < 0.001$ ]. Moreover, patients with 5 or 297  
 263 more SDI score had significantly shortened mean survival 298  
 264 time than patients with 4 or less SDI score [24.05 years 299  
 265 (95% CI, 20.75–27.35) vs. 43.79 years (42.66–44.93), respec- 300  
 266 tively,  $p < 0.0001$ ] (Fig. 4a, b).

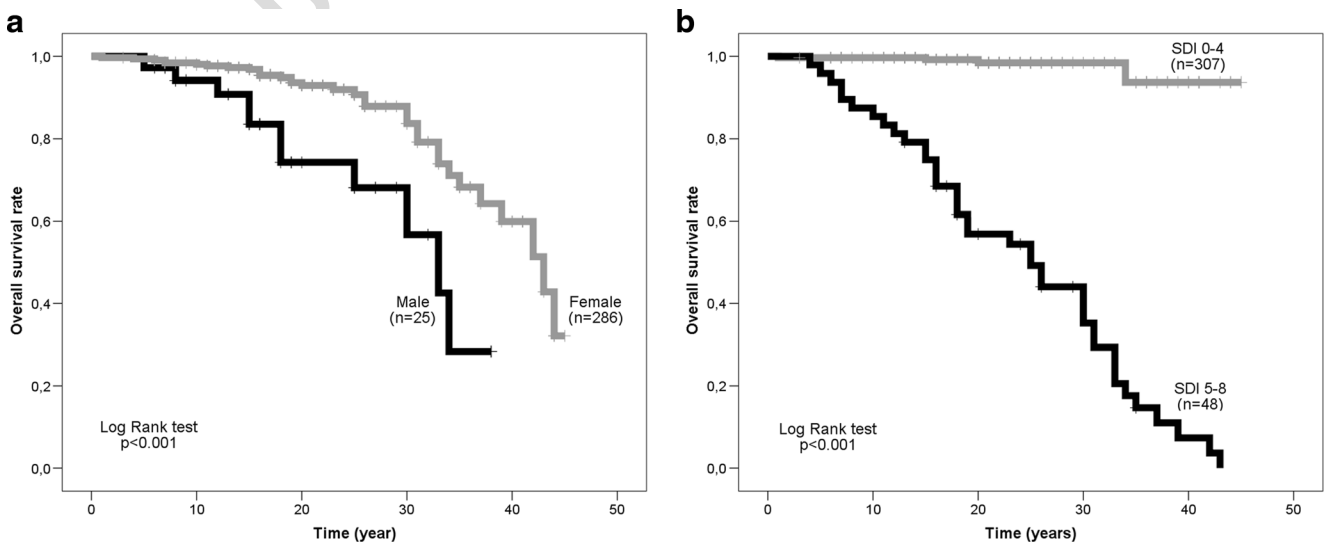
Cox regression analyses revealed three independent prog- 267  
 nostic factors: male gender, >4 SDI score and higher cumula- 268  
 tive glucocorticoid doses have significant negative effect on 269  
 disease outcome [male gender: hazard ratio (HR), 2.785 (95% 270  
 CI, 1.35–5.719), respectively,  $p = 0.005$ ; >4 SDI score: HR, 271  
 55.12 (95% CI, 19.15–158.63), respectively,  $p < 0.001$ ; cumu- 272  
 lative glucocorticoid doses: HR, 1.02 (95% CI, 1.006– 273  
 1.035), respectively,  $p = 0.005$ ]. 274

**Discussion**

In SLE, chronic organ damage has become an increasingly im- 276  
 portant factor beyond disease activity. Many factors such as 277  
 geographic and ethnic determinants can affect the severity and 278  
 course of the disease as well as the development of organ dam- 279  
 age. In spite of the wealth in international data, our information 280  
 on chronic organ damage and understanding of its determinants 281  
 in SLE patients in East-Central Europe is incomplete, and the 282  
 results measured by various centres diverge on several points. 283

Our results show that the patient's gender does not influ- 284  
 ence the development of chronic organ damages. Yee et al. 285  
 and Estevez del Toro et al. obtained similar results in British 286  
 and Cuban patients, respectively [17, 18]. In contrast, 287  
 Andrade et al. found that male patients developed chronic 288  
 organ damage faster and in larger numbers [19]. The incidence 289  
 of the most common damages can vary. Among our patients, 290  
 the most frequent damages were found in the cardiovascular 291  
 and neuropsychiatric organ systems. The largest numbers of 292  
 chronic organ damage were found in the renal and musculo- 293  
 skeletal systems [20], the musculoskeletal and dermal systems 294  
 [18] and the neuropsychiatric system [21]. 295

We made the assumption that among patients with longer 296  
 disease duration, the number of chronic organ damages may 297



**Fig. 4** Kaplan-Meier survival plots for patients subgroups. **a** Male and female patients. **b** Patients with SDI value >4 and <5

298 be increased. An additional complicating factor was that these  
 299 patients might have been treated with several types of immu-  
 300 nosuppressive therapies. Duration of disease has been desig-  
 301 nated as a factor in chronic damage by several centres [21].  
 302 There is disagreement in the results as to whether SDI value  
 303 shows a linear increase with disease duration. A gradual in-  
 304 crease was found by Cassano [22] in the Argentinian SLE  
 305 population, and a similar linear increase was measured by  
 306 Gladman et al. [23]. In agreement with our results, a gradual  
 307 increase followed by a “plateau phase” after certain duration  
 308 of disease was described by Becker-Merock and Nossent [24].  
 309 Interestingly, we found that the prevalence of chronic damage  
 310 was 77.9% in our Hungarian SLE cohort, which is higher  
 311 compared with other European cohort [5, 17]. This difference  
 312 can be explained by our results, since the follow-up of our  
 313 patients was longer, compared with the other cohorts, and  
 314 based on our observations, a significant increase in SDI values  
 315 develops typically 10 years after diagnosis.

316 During the course of SLE, chronic damage may develop with  
 317 a higher frequency among patients with increased disease activ-  
 318 ity. As described earlier by Lopez et al., disease activity mea-  
 319 sured by BILAG predicted later damages [25]. In their 5-year  
 320 prospective study, Stoll et al. found that disease activity defined  
 321 the development of chronic damages [26]. Although we did not  
 322 detect a significant difference in the course of the present study,  
 323 the number of patients showing active disease during the prior  
 324 10 years was higher among SLE patients with higher SDI values.

325 Similar to our results, Maddison et al. described the  
 326 role of mean age at the time of diagnosis. Higher SDI  
 327 values were found among patients who were diagnosed  
 328 after the age of 40 years than those diagnosed under 40  
 329 [27]. In contrast, Morgan et al. found that young and  
 330 adolescent SLE patients sustain more damage over time  
 331 [28]. In his study of Chinese lupus patients, Feng com-  
 332 pared damages in patients with SLE diagnosed in child-  
 333 hood (under 18 years of age), youth (between 18 and  
 334 45 years of age) and old age (above 45 years of age);  
 335 no difference was found in the damage indexes [29].

336 Various aspects of the effects of glucocorticoids on chronic  
 337 organ damage were evaluated. Some publications examined  
 338 cumulative doses of glucocorticoids [30], while others studied  
 339 the average daily doses [20] or the potential effect of paren-  
 340 teral glucocorticoid therapy [31]. Mae Thaner et al. found that  
 341 the risk of irreversible damage increased with an increase of  
 342 the glucocorticoid dose. However, there was no significant  
 343 difference in the development of damage with administration  
 344 of low-dose (<180 mg/month) prednisolone [30]. Gladmann  
 345 et al. found that the amount of glucocorticoid administered  
 346 had an unequivocal effect on the development of cataracts  
 347 and a likely effect on cardiovascular events [23]. We also  
 348 found a strong association with high-dose glucocorticoid ther-  
 349 apy cataract and osteoporosis. Cumulative glucocorticoid  
 350 dose influenced also the cerebrovascular events, myopathy,

lower extremity claudication and avascular necrosis of the  
 femoral head, but the difference was not significant.

Regarding immunosuppressive agents, we described the  
 beneficial effect of chloroquine. Data from the Lumina cohort  
 found that the SDI values of patients given initial chloroquine  
 therapy were lower [32]. According to Akhavan et al., in the  
 case of patients treated with chloroquine, less damage could  
 be expected during the 3 years after diagnosis [33].

Several other groups described that SLE patients treated  
 with cyclophosphamide had higher mean SDI values [20,  
 34]. However, we did not detect any direct correlation be-  
 tween this and other immunosuppressive agents and the fre-  
 quency of chronic organ damage among our patients. In con-  
 trast, Mok and Akhavan described a significant correlation  
 between azathioprine and chronic damage in Chinese and  
 Canadian patients with SLE [33, 35]. A recent study demon-  
 strated the possible role of anti-phospholipid antibodies in the  
 development of organ damage [36]. We did not reveal any  
 associations between serological features and SDI; however,  
 the more careful assessment of anti-phospholipid antibody-  
 positive patients is undoubtedly necessary.

Significant gender differences were found in survival ratios;  
 moreover, elevated SDI scores and higher cumulative doses of  
 glucocorticoids increased mortality risk. This is in accordance  
 with the fact that mortality ratios can improve and toxic adverse  
 effects of glucocorticoids can be decreased by the usage of  
 newer drugs with reduced glucocorticoid doses [37].

Our results demonstrate that as long-term survival in SLE  
 improves, it becomes increasingly important to survey the re-  
 sults and to identify the determinants of chronic organ damage.  
 Our data confirmed that disease duration, age at diagnosis and  
 chronic high-dose glucocorticoid therapy have significant ef-  
 fects on the development of chronic organ damage in the  
 Hungarian patients with SLE. Our data are representative of  
 East-Central European SLE population as well. Additionally,  
 we confirmed the protective effect of chloroquine. Most of the  
 chronic organ damage occurs in the cardiovascular and the  
 neuropsychiatric systems; thus, regular follow-up, screening  
 and adequate therapy are essential for the best clinical outcome.

**Compliance with ethical standards** All experiments carried out in the  
 study were in compliance with the Declaration of Helsinki.

**Disclosures** None.

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AUTHOR QUERY

**AUTHOR PLEASE ANSWER QUERY.**

- Q1. “Systemic Lupus Erythematosus Disease Activity Index” was provided as the definition for “SLEDAI.” Please check and amend as necessary.

UNCORRECTED PROOF