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Complement levels during the first trimester predict disease flare and adverse pregnancy outcomes in Systemic Lupus Erythematosus: a network meta-analysis on 532 pregnancies

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Short Title: Complement levels predict flares in lupus pregnancies

Key words: SLE; Systemic Lupus Erythematosus; Pregnancy; Complement

Abstract

Background: Complement levels have been proposed as candidate biomarkers of disease activity and obstetric risk in lupus pregnancies, but their reliability has been questioned due to the physiologic fluctuations of complement levels during gestation. Thus, this network meta-analysis aimed at assessing the clinical significance of complement fluctuations in lupus pregnant women.

Methods: Corresponding authors of 19 studies meeting inclusion criteria were invited to contribute with additional data including C3 and C4 levels [before pregnancy, at conception, in every trimester (T) and 3 months after delivery]; data were pooled together in a network meta-analysis.

Results: A total of 532 lupus women from four studies were included in the analysis. In SLE women, C3 and C4 increased progressively our ng Bestation: levels remained stable during T1 and peaked in T2 to decrease in T3. Patients with previous lupus nephritis (LN) and those who experienced flares during pregnancy had significantly lower mean levels of C3 and C4 at all timepoints. The lowest levels of complement were observed, particularly during T1, in patients with LN and gestational flare. Poth reduction and the lack of increase of C3 and C4 levels at T1 versus conception were as occated with gestational flares, particularly in LN patients. Pregnancies with flare had a statistically significant higher rate of maternal and fetal complications (60%vs.50.3%; p=0.03).

Conclusions: Low complement levels, particularly in T1, were associated with a higher frequency of gestational flare. Either reduction or smaller increase of C3 and/or C4 levels, even within normal range, might predict flares especially in early gestation.



Highlights:

- C3/C4 values progressively increase in lupus pregnancy as in physiological pregnancy, but the peak is in the second trimester, then there is a decline, particularly in C3.
- Patients with flare during pregnancy and those with previous LN showed significantly lower complement levels than those who did not have flare and did not have renal involvement, respectively.
- The lowest complement levels pre-pregnancy and at conception were observed in patients with previous LN and related gestational flare.
- The fluctuations of C3 and C4 levels in the first trin. ester versus conception displayed the highest clinical significance in predicting both SI Energy and APO as compared to variations between other timepoints.
- A decrease, stable values, or an increase smaller than 5 mg/dL in C3 levels between conception and the first trimester ware all factors associated with an increase in the risk of SLE flares and APO.
- Validation studies will clarify these findings and define the role of C3 and C4 levels during the preconception period/early pregnancy as tools for predicting SLE flares and APO and implementing individualized treatment strategies.
- In lupus patients, C3 and C4 monitoring should be incorporated into the clinical assessment before conception and throughout the entire pregnancy.

1. Introduction

Systemic lupus erythematosus (SLE) is a prototypical immune complex-mediated disease characterized by a wide spectrum of disease phenotypes with heterogeneous courses and progression, varying from persistently low, relapsing-remitting, to persistently high disease activity [1]. The epidemiology of SLE, which mainly presents in young women of childbearing age [2], accounts for the fact that clinicians assist lupus patients very often in their journey towards motherhood. To explain such epidemiological female predominance, several hypotheses have been formulated: candidate risk genes for SLE map on the X caroniosome, and estrogens favour autoimmunity by promoting B-cell maturation, antibody crochiction, Th2 responses, and survival of autoreactive cells [3].

As expected, pregnancy can impact SLE disease activity, and in turn SLE may affect obstetric outcomes. Pregnancy in women with SLE hat always been regarded as at high risk; however, the significant advancements made in the coral disease management have led to a net improvement of both maternal and fetal outcomes [4,5]. Nevertheless, pregnancy still represents a challenge in women with SLE, especially in those with renal involvement, due to the hazard of disease flare, gestational diabetes and place ta-related disorders including pre-eclampsia (PE), as well as fetal complications such as micropriages, fetal loss, intrauterine growth restriction, prematurity, and neonatal lupus [6]. Reliable biomarkers to stratify the risk of a disease flare during pregnancy and to early detect adverse pregnancy outcomes (APO) in pregnant lupus women are still lacking. Complement levels have been proposed as candidate biomarkers of disease activity and of obstetric risk in lupus pregnancies, but their reliability has been questioned due to the physiologic fluctuation of complement levels during gestation [7,8]. Recently, one cohort study described a predictive role of low pre-pregnancy C4 levels towards disease flare during pregnancy [9], while another one found that low-pregnancy C3 levels were associated with preterm birth [10].

In order to optimize the interpretation of available data on the variation of complement levels during SLE pregnancy, we performed a network meta-analysis to assess the fluctuations of C3 and C4 levels from preconception period, throughout pregnancy, and up to 3 months after delivery and to evaluate the association of complement levels with the occurrence of disease flares and/or APO.

2. Patients Methods

2.1 Systematic literature review

A detailed literature search strategy has been developed *a priori* to identify articles that reported findings from available prospective studies investigating pregnancies in patients with SLE from January 2002 to December 2020. Key words and subject terms included: (("longitudinal studies"[MeSH Terms] OR ("longitudinal"[All Fields] AND "studies"[All Fields]) OR "longitudinal studies"[All Fields] OR "prospective"[All Fields] OR "prospective; [All Fields]) AND ("lupus vulgaris"[MeSH Terms] OR ("lupus"[All Fields] AND "vulgar s"[**, "Fields]) OR "lupus vulgaris"[All Fields] OR "lupus erythematosus, "stemic"[MeSH Terms] OR ("lupus"[All Fields] AND "systemic"[All Fields]) OR "systemic lupus erythematosus"[All Fields]) AND "systemic"[All Fields]) OR "pregnancy"[All Fields] OR "pregnancy"[All Fields]) OR "pregnancy"[All Fields]).

The search strategy was applied to Cv d MEDLINE, In-Process and Other Non-Indexed Citation from January 2002 to December 2020. Figure 1 resumes the search strategy.

Retrieved papers were further screened upon additional inclusion criteria in order to refine the search strategy. Inclusion cr teria included: a) prospective design, b) a sample size of at least 50 lupus patients, c) exclusion of miscarriages before 12 weeks of gestation as obstetric outcome.

Given the nature of this study, ethics committee approval was not required.

2.2 Data collection and analysis

Two review Authors (M.R. and I.C.) independently assessed studies for inclusion. One review Author completed data extraction, which was checked by a second review Author. A total of 19 studies were finally selected for data request. Each corresponding Author of the selected manuscripts was invited to contribute with additional data of individual pregnancies that were not presented in the published manuscript, including complement levels, C3 and C4 separately, at 6 months before pregnancy, at conception, during the first trinnster (T1), during the second trimester (T2), during the third trimester (T3), and 3 months after delivery (post-partum, PP). Further details on the number of pregnancies, patients classification, diagnosis at conception, treatment during pregnancy, occurrence of flares during restation, as well as maternal and fetal outcomes were also recorded. We performed a network meta-analysis within a Bayesian framework as previously described [11].

2.3 Statistical analysis

Categorical variables are presented as numbers (%) and continuous variables are expressed as mean ± standard deviation SD). The significance of baseline differences was determined by the chi-squared test, Fisher's exact test or the unpaired t-test, as appropriate. Correlation analysis, linear regression, and Odds Ratio (OR) were also performed. Missing data were approached with mean substitution system. A two-sided P-value <0.05 was considered statistically significant. All statistical analyses were performed using SPSS version 26.0 (IBM, Armonk, NY, USA).

2.4 Study variables definitions

SLE, lupus nephritis (LN), and antiphospholipid syndrome (APS) diagnosis and classification were based upon each study definition [12–14]. SLE flare was defined by the need of new immunosuppressive therapy or increase in the dosage of prednisone ≥10 mg/day.

APO were defined as follows:

- a) fetal death after 12 weeks' gestation unexplained by chron osomal abnormalities, anatomic malformation, or congenital infection;
- b) neonatal death before hospital discharge due to consolications related to prematurity or placental insufficiency (e.g., abnormal fetal surveillance test results, abnormal Doppler flow velocimetry waveform analysis suggestive of fetal homoxemia, or oligohydramnios, or both);
- c) preterm delivery or pregnancy loss at less tion 36 weeks due to gestational hypertension, PE, or placental insufficiency;
- d) small-for gestational-age neon ite, defined as one with a birth weight below the 5th percentile without anatomical or chromosemal abnormalities.

The fluctuation of C3 and C2 levels between T1 and conception was defined as Δ C3_{T1-conception} and Δ C4_{T1-conception}. When the decrease in C3 levels between T1 and conception was below 2 mg/dl or the increase in C3 at T1 *versus* conception was below 4 mg/dl (defined using two standard deviations from mean, as per Westgard rules), Δ C3_{T1-conception} was considered as clinically not relevant.

3. Results

3.1 C3 and C4 levels progressively increased during gestation in women with SLE

A total of 532 SLE women from 4 studies were included in the analysis [15–18]. APS had been diagnosed in 68 women (12.8%), while 82 patients (15.4%) were positive for antiphospholipid antibodies (aPL) without overt clinical manifestations of APS (referred as "aPL carriers"). As detailed in Table 1 and visually presented in Figure 2A, both C3 and C4 levels increased progressively in women with SLE during gestation. In particular C2 and C4 levels remained stable during T1 and peaked at T2, then decreased during T3. Af 3 months after delivery, a different behavior was noted for C3 and C4: C3 continued to decrease whereas C4 levels in the PP period were higher than those registered in T3.

3.2 Patients with flares during pregnant, displayed significantly lower levels of complement compared to patients without gestational flare

A flare during pregnancy was observed in 170 patients (32%). Levels of both C3 and C4 were lower at all timepoints in subjects who experienced flares during pregnancy (C3 at T1 78.3 \pm 22.8 *versus* 100.5 \pm 20.7, p<0.001; C3 at T2 94.2 \pm 13.4 *versus* 115.7 \pm 12.3, p<0.001; C3 at T3 99 \pm 18.6 *versus* 111.4 \pm 16, p<0.001; C3 at PP 92.4 \pm 15.7 *versus* 102.6 \pm 13.4, p<0.001; Table 1 and Figure 2B).

The physiological increase in complement levels throughout gestation was rather marked among patients who did not experience a disease flare while pregnant. Complete data on complement levels fluctuation at all time-points in patients experiencing a gestational flare vs. those who did not present a disease flare while pregnant are listed in Table 1 and illustrated in Figure 2B.

3.3 Patients with LN displayed significantly lower levels of complement compared to patients without renal involvement

LN had been diagnosed in 237 women (44.5%). Patients with LN had significantly lower levels of complement when compared to patients without renal involvement (C3 at conception 96.1 \pm 13.9 versus 91.1 \pm 13, p<0.001; C3 at T1 84.6 \pm 32.2 versus 98.4 \pm 14.1, p<0.001; C3 at PP 93.4 \pm 12 versus 103.1 \pm 15.4, p<0.001; C4 at conception 15.4 \pm 4.1 versus 13.9 \pm 2.8, p<0.001; C4 at T1 15 \pm 7.8 versus 16.3 \pm 2.8, p<0.001; C4 at PP 16.2 \pm 4.3 versus 19.8 \pm 6.9, p<0.001, Table 1and Figure 2C).

3.4 Patients with previous LN and flare during pregnancy displayed the lowest complement levels

A flare during pregnancy was observed in 73 women with a previous diagnosis of LN. The lowest levels of complement, both for C3 and C4, were observed in patients with a previous diagnosis of LN who experienced a flare during pregnancy. Complete data are listed in Table 1 and visually represented in Figure 2C.

3.5 The fluctuations of C3 and C4 levels at T1 *versus* conception displayed the highest clinical significance in predicting disease flares

When analyzing the fluctuations of complement levels between different timepoints, the variations in both C3 and C4 between levels assessed at T1 *versus* those recorded at conception emerged as the most clinically significant. Indeed, the differential values in both C3 and C4at T1 *versus* at conception (defined as Δ C3_{T1-conception} and Δ C4_{T1-conception}, respectively) were significantly

lower in patients with LN when compared to patients without renal involvement (Δ C3 0.5±53 versus16.6±34.3, p<0.001; Δ C4 1.5±9.1 versus4.5 ±6.3, p<0.001).

Women who experienced a flare during pregnancy had lower $\Delta C3_{T1-conception}$ and $\Delta C4_{T1-conception}$ ($\Delta C3_{T1-conception}$ -6.7±48.8 *versus* 18.8±37.6, p<0.001; $\Delta C4$ 1.2±8.1 *versus* 4.4±7.1, p <0.001). The lowest levels of $\Delta C3_{T1-conception}$ and $\Delta C4_{T1-conception}$ were reported in patients that were diagnosed with LN and experienced flares during pregnancy ($\Delta C3_{T1-conception}$ -36.1±42.6; $\Delta C_{T1-conception}$ -1.1±8.5).

A decrease in Δ C3_{T1-conception} yielded an OR for flare during p regrancy of 3.1 (CI 95% 2.1-4.8) when below 5 mg/dL, an OR that increased up to 3.9 (CI 95% 2.5-5) when below 15 mg/dL.

Similar figures emerged when assessing the association between $\Delta C3_{T1-conception}$ and a prior diagnosis of LN: $\Delta C3_{T1-conception} \leq 5$ mg/dL crank yea an OR for a prior diagnosis of LN of 6.1 (CI 95% 3.9-9.6) while $\Delta C3_{T1-conception} \leq 10$ mg/dL conveyed an OR of 7.2 (CI 95% 4.5-11.7). Interestingly, even the lack of clinically relevant changer in the complement levels between T1 and conception was associated with both previou. LN diagnosis (OR 2.2; CI 95% 1.3-3.6) and development of flare during pregnancy (OR 5.2; CI 95% 2.9-9.3). Table 2 resumes the results of the coefficient of risk conveyed by different $\Delta C3_{1-conception}$ levels upon LN diagnosis or presence of flare.

3.6 The fluctuations of C3 and C4 levels at T1 *versus* conception displayed the highest clinical significance in predicting APO

Preterm delivery or miscarriage at less than 36 weeks were more frequent in women with a previous diagnosis of APS (39.7% *versus* 23%; p=0.003), in patients who developed flares during pregnancy irrespectively of a concomitant diagnosis of LN (42.5% *versus* 28%; p= 0.01 in patients

with LN and 34% *versus* 17.2%; p= 0.01 in those without a diagnosis of LN). Additionally, fetal death was more frequent in patients with a diagnosis of LN and positive aPL (4/30 *versus* 6/206; p=0.008).

When computing all APO together, higher rates of complications were reported in patients with a previous diagnosis of APS (88.2% *versus* 56%; p< 0.0001) as well as LN (67.9% *versus* 53.9%; p<0.0001) and occurrence of flare during pregnancy (91.2% *versus* 45.6%; p<0.0001).

 Δ C3_{T1-conception} \leq 5mg/dL and no changes of Δ C3_{T1-conception} were \sim th associated with higher rates of overall APO (63.4% *versus* 45.6%; p=0.003 and 58.5% *vers is*7 \geq 8%; p=0.02, respectively).

4. Discussion

The present network meta-analysis, which includes more than 500 pregnant lupus patients from 4 international independent studies, allowed us to clearly assess the clinical relevance of complement monitoring during gestation to predict both disease flares and APO. Levels of C3 and C4 emerged as a reliable biomarker to identify those women who are at higher risk of developing disease flares and APO, even in case of a concomitant diagnosis of LN [19]. These findings are extremely relevant from a clinical perspective given that, despite the substantial improvements accomplished in the management of SLE patients, 50% of lunus women might develop a flare during gestation, with severe organ involvement occurring in up to 25% of cases [20,21]. Unfortunately, the current lack of reliable biomarkers and validated tools for the assessment of disease activity during pregnancy limits our ability to predict which subjects will experience disease worsening and/or APO. In the lant new decades, several scoring systems have been developed to assess lupus activity and the . isk of flare during pregnancy. Most of these tools, such as the LAI in Pregnancy (LAI-P), the See-Pregnancy Disease Activity Index (SLEPDAI), and the modified SLAM3 (m-SLAM) [22] include hypocomplementemia (C3 and C4). These clinimetric instruments have been riea ed by modifying existing lupus activity indexes in order to differentiate between disease-specific features and physiologic changes occurring during gestation. Although promising, these pregnancy-adapted scores have not been extensively validated in large prospective cohorts and therefore their current employment in clinical practice is strongly limited. Similarly, C3 and C4 levels should be carefully evaluated in pregnant lupus women as complement serum levels rise throughout the course of normal gestation [23]. This study confirms that complement levels fluctuate over the gestational course also in SLE women: values of C3 and C4 remained stable at early stages of pregnancy, then progressively increased during the second trimester of gestation; once reached the highest levels, both C3 and C4 showed

a decline with discrepant behaviors after delivery, resulting in a constant rise of C4 values and a progressive decrease of C3. Interestingly, we observed that lupus patients who experienced a clinical flare during pregnancy had significantly lower mean values of C3 and C4 throughout the entire gestation compared with patients with stable disease activity. If our data confirm the relevance of complement as a monitoring tool of lupus disease activity even during gestation, it should be mentioned that the consensus about the reliability of complement in predicting SLE flare is not unanimous. Indeed, its relevance has been questioned by few studies [24–27], most likely due to the methodological challenges of accurately meast ring circulating complement levels as well as to the inappropriate designs of clinical studies [28]. Nevertheless, despite these inconsistencies, it is universally accepted that complyment activation in SLE is mirrored by a secondary decline of circulating complement levels and a parallel increase in complement split products and circulating levels of complement processins (C3 and C4) are extensively used in clinical practice for classification and diagnostic purposes, monitoring of disease activity and follow-up. Similarly, the clinical significance of lov (3 and C4 circulating levels as biomarker for LN is still matter of research [29]. Whereas a significant drop in C4 levels can be observed even two months prior to renal flare occurrence, a recline in C3 was shown to be influenced by genetic variants of factor H, which regulates C?-convertase in the alternative pathway [30]. In addition, elevated titers of autoantibodies directed against C1q have been described as better predictors of renal involvement in SLE patients compared to C3 and C4, although with inconclusive results [30]. Further analysis of our data revealed significantly lower levels of C3 and C4 in pregnant patients with LN at all time-points considered, from conception throughout pregnancy and until 3 months following delivery, consistently with what reported by other authors [30]. Most importantly, the present study also highlights that those patients with both previous LN and disease flare during gestation had the lowest complement levels, suggesting that levels of C3 and C4 below the normal

threshold before conception can serve as predictor of flare during pregnancy in this high-risk group of patients [31].

To better evaluate the fluctuations of C3 and C4 by minimizing the confounding effect of cut-off variability and inter-assay heterogeneity among the four different cohorts, as well as the potential influence of genetic variants, the analysis in this network meta-analysis also focused on the differential levels of circulating C3 and C4 values (ΔC3 and ΔC4) between different trimesters of gestation, rather than the mere absolute levels or the dictatomous categorization into hypocomplementemia versus normocomplementemia. This approach allowed us to determine that the most informative data in clinical practice consists in the lack of physiological increase in C3 and C4 values in the first trimester of gestation as compared to conception: women who experienced a lupus flare during gestation displayed the lowest Δ C3 and Δ C4 during the first trimester versus at conception. In addition, "le less pronounced the increase in C3 levels is from conception throughout the first trimestor of gestation, the higher the risk of developing disease flare with an OR up to 3.9 when ΔC_3 is below 15 mg/dl. The same conclusion can be extrapolated to pregnant women with renal involvement and the occurrence of flare during gestation, a subset of patients in which a smal ΔC carried an even higher risk of disease flare (OR 5.2). Despite the significance of C4 variations during pregnancy in predicting both APO and disease flare, we decided to emphasize the results obtained when focusing on C3 variations. In fact, from a practical point of view, and based on the more extended range of C3 values, ΔC3 might be easier to assess and more informative for clinicians.

The data gathered in this meta-analysis allowed us to investigate also the role of complement levels in predicting obstetric morbidity among lupus women. Women with lower levels of both C3 and C4 prior to conception and during the entire gestation are more likely to experience APO: a

 Δ C3 below 5 mg/dl between the first trimester and at conception as well as no changes in Δ C3 at these time-points were associated with an overall higher rate of APO. These findings are consistent with the available literature, which traditionally enlists hypocomplementemia, together with active LN at conception, previous history of LN, aPL positivity and high disease activity before conception, as major determinants of poor maternal and fetal outcomes in lupus women [6]. The relationship between complement levels and APO should not be surprising, given the multifaceted role of the complement cascade in pregnant lupus women. On or e hand, the complement system, with more than 30 plasma proteins and receptors, represents a key element of the innate immunity response that contributes to the progression of SLE through the stimulation of inflammation and the removal of immune complexes, calls, and apoptotic debris [32]. Importantly, SLE onset, disease activity and organ damage have been all linked to complement activation and consumption, as well as to complement deficiencies [33]. On the other hand, a consistent stream of data has progressively shown that the complement cascade exerts a pivotal role throughout all the stages of physiologic gestation (concept or, embryo implantation, placentation, fetal growth, and labor) and the fine tuning of the expression of complement factors, receptors and inhibitors during gestation, with their increased be atic synthesis, is mandatory to ensure a successful pregnancy [34].

This study presents some limitations that should be acknowledged. First, the limited number of included studies does not encompass the whole prospective experience in lupus pregnancy available in the literature. Second, the geopolitical representation of the included cohorts does not comprehend North America, Asia-Pacific and Africa, thus reducing the generalizability of our conclusions. Third, since SLE is an extremely heterogeneous condition, the inclusion of patients with distinct clinical profile might limit the reproducibility of the observed results. Fourth, given the nature of the study, the lack of a control group (e.g. healthy subjects) represents another

limitation. Despite the acknowledged limitations, our study has indeed some strengths: the high number of included patients, the prospective design of the considered studies, and lupus diagnosis assessed with homogeneous criteria across different cohorts [12–14]. Moreover, despite the absence of complement levels adjustment for gestational state or trimester [35], cut-off values for circulating levels C3 and C4 were comparable among different cohorts.

This network meta-analysis suggests the role of C3 and C4 levels/fluctuations before conception and in early pregnancy as predictors of SLE flares and APO later in the pregnancy course. Particularly, the lack of increase in C3 and C4 levels during another thirteen weeks of gestation appeared as a strong predictor of flare, especially in women with previous LN. These findings deserve further validation in order to define the role of complement as a biomarker that can inform risk stratification and guide individualized treatment decisions in women with SLE who are pregnant or planning to get pregnant.

5. References:

- 1. Aringer M, Johnson SR. Classifying and diagnosing systemic lupus erythematosus in the 21st century. Rheumatology (Oxford) 2020;59:V4–11.
- 2. Tian J, Zhang D, Yao X, Huang Y, Lu Q. Global epidemiology of systemic lupus erythematosus: a comprehensive systematic analysis and modelling study. Ann Rheum Dis 2022;ard-2022-223035.
- 3. Dao KH, Bermas BL. Systemic Lupus Erythematosus Management in Pregnancy. Int J Womens Health 2022;14:199–211.
- 4. Buyon JP, Kim MY, Guerra MM, Laskin CA, Petri M, Lockshin MD, et al. Predictors of Pregnancy Outcomes in Patients With Lupus: A Cohort Study. Ann Intern Med 2015;163:153–63.
- 5. Andreoli L, Gerardi MC, Fernandes M, Bortoluzzi A, Bellando-Racciono S, Brucato A, et al. Disease activity assessment of rheumatic diseases during pregnancy: a comprehensive review of indices used in clinical studies. Autoimmun Rev 2019;18:164–76.
- 6. Petri M. Pregnancy and Systemic Lupus Erythematosus Res. Pract Res Clin Obstet Gynaecol 2020;64:24–30.
- 7. Chakravarty EF, Colón I, Langen ES, Nix DA, El-Saye 'Y', Genovese MC, et al. Factors that predict prematurity and preeclampsia in pregnancies has are complicated by systemic lupus erythematosus. Am J Obstet Gynecol 20°5;1)2:1097–904.
- 8. Saleh M, Compagno M, Pihl S, Strevens H, Pelsson B, Wetterö J, et al. Variation of Complement Protein Levels in Maternal Plasma and Umbilical Cord Blood during Normal Pregnancy: An Observational Study. J Clin Med 20.12 11.
- 9. Crisafulli F, Andreoli L, Zucchi D, Peggia R, Gerardi MC, Lini D, et al. Variations of C3 and C4 Before and During Pregnancy in Systemic Lupus Erythematosus: Association With Disease Flares and Obstetric Outcomes. J Rhematol [Internet] 2023 [cited 2023 Jul 12]; jrheum. 2022-1135. Available from: https://pubmed.icb.nlm.nih.gov/37127323/
- 10. Hiramatsu Y, Isoda K, Kutani T, Nakamura E, Wada Y, Fujiki Y, et al. Pre-pregnancy serum complement C3 level is a predictor of preterm birth for pregnancies with systemic lupus erythematosus. Arthritis Res Ther [Internet] 2021 [cited 2023 Jul 12];23. Available from: https://pubmed.ncbi.nlm.nih.gov/33980284/
- 11. N Nyaga V, Arbyn M, Aerts M. Beta-binomial analysis of variance model for network meta-analysis of diagnostic test accuracy data. Stat Methods Med Res 2018;27:2554–66.
- 12. Hochberg MC. Updating the American College of Rheumatology revised criteria for the classification of systemic lupus erythematosus. Arthritis Rheum 1997;40:1725.
- 13. Miyakis S, Lockshin MD, Atsumi T, Branch DW, Brey RL, Cervera R, et al. International consensus statement on an update of the classification criteria for definite antiphospholipid syndrome (APS). J Thromb Haemost 2006;4:295–306.

- 14. Tan EM, Cohen AS, Fries JF, Masi AT, McShane DJ, Rothfield NF, et al. The 1982 revised criteria for the classification of systemic lupus erythematosus. Arthritis Rheum 1982;25:1271–7.
- 15. Moroni G, Doria A, Giglio E et al. Maternal outcome in pregnant women with lupus nephritis. A prospective multicenter study. J Autoimmun. 2016;74:194-200.
- 16. Borella E, Lojacono A, Gatto M, Andreoli L, Taglietti M, Iaccarino L, et al. Predictors of maternal and fetal complications in SLE patients: a prospective study. Immunol Res 2014;60:170–6.
- 17. Saavedra MÁ, Miranda-Hernández D, Lara-Mejía A, Sánchez A, Morales S, Cruz-Reyes C, et al. Use of antimalarial drugs is associated with a lower risk of preeclampsia in lupus pregnancy: A prospective cohort study. Int J Rheum Dis 2020;23:633–40.
- 18. Rodrigues BC, Lacerda MI, Ramires de Jesús GR, Cunha dos Sant is F, Ramires de Jesús N, Levy RA, et al. The impact of different classes of lupus nephritis on maternal and fetal outcomes: a cohort study of 147 pregnancies. Lupus 2019;28:492–500.
- 19. Weinstein A, Alexander R V., Zack DJ. A Review of Complement Activation in SLE. Curr Rheumatol Rep 2021;23:4–11.
- 20. Buyon JP, Kim MY, Guerra MM, Lu S, Reeves E, Petri M, Cal. Kidney outcomes and risk factors for nephritis (flare/de novo) in a multiethnic cohort of pregnant patients with lupus. Clinical Journal of the American Society of Nephrology 2017;12: 40 C.
- 21. Larosa M, Le Guern V, Guettrot-Imbert Morel N, Abisror N, Morati-Hafsaoui C, et al. Evaluation of lupus anticoagulant, damage, and remission as predictors of pregnancy complications in systemic lupus erythematosus: the French GR? surdy. Rheumatology (Oxford) 2022;61:3657–66.
- 22. Buyon JP, Kalunian KC, Ramsey-Goldman R, Petri MA, Lockshin MD, Ruiz-Irastorza G, et al. Assessing disease activity in SLE patients using pregnancy. Lupus 1999;8:677–84.
- 23. Kim MY, Guerra MM, Kaplowicz E, Laskin CA, Petri M, Branch DW, et al. Complement activation predicts adverse pregnancy cuccome in patients with systemic lupus erythematosus and/or antiphospholipid a: tib. dier. Ann Rheum Dis 2018;77:549–55.
- 24. Esdaile JM, Abrahamcu icz M, Joseph L, MacKenzie T, Li Y, Danoff D. Laboratory tests as predictors of disease exacerbations in systemic lupus erythematosus. Why some tests fail. Arthritis Rheum 1996;39:370–8.
- 25. Merrill JT, Petri MA, Buyon J, Ramsey-Goldman R, Kalunian K, Putterman C, et al. Erythrocyte-bound C4d in combination with complement and autoantibody status for the monitoring of SLE. Lupus Sci Med 2018;5.
- 26. Steiman AJ, Gladman DD, Ibañez D, Urowitz MB. Prolonged serologically active clinically quiescent systemic lupus erythematosus: frequency and outcome. J Rheumatol 2010;37:1822–7.
- 27. Sandhu V, Quan M. SLE and Serum Complement: Causative, Concomitant or Coincidental? Open Rheumatol J 2017;11:113–22.
- 28. Liu CC, Manzi S, Danchenko N, Ahearn JM. New advances in measurement of complement activation: lessons of systemic lupus erythematosus. Curr Rheumatol Rep 2004;6:375–81.

- 29. Birmingham DJ, Irshaid F, Nagaraja HN, Zou X, Tsao BP, Wu H, et al. The complex nature of serum C3 and C4 as biomarkers of lupus renal flare. Lupus 2010;19:1272–80.
- 30. Mok CC. Epidemiology and survival of systemic lupus erythematosus in Hong Kong Chinese. Lupus 2011;20:767–71.
- 31. Swaak AJG, Groenwold J, Bronsveld W. Predictive value of complement profiles and anti-dsDNA in systemic lupus erythematosus. Ann Rheum Dis 1986;45:359–66.
- 32. Walport MJ. Complement. First of two parts. N Engl J Med 2001;344:1058–66.
- 33. Chighizola CB, Lonati PA, Trespidi L, Meroni PL, Tedesco F. The Complement System in the Pathophysiology of Pregnancy and in Systemic Autoimmune Rheumatic Diseases During Pregnancy. Front Immunol 2020;11:1–11.
- 34. Girardi G, Lingo JJ, Fleming SD, Regal JF. Essential Role of Complement in Pregnancy: From Implantation to Parturition and Beyond. Front Immunol 2020, 1:1-17.
- 35. Buyon JP, Kim M GM et al. Predictors of Pregnancy Outco. e in a Prospective, Multiethnic Cohort of Lupus Patients. Annals of Internal Medicine 2015;163 153 -63.

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Legends of Figures and Tables

Figure 1. Literature search strategy.

Figure 2. Complement levels fluctuations over 6 time points (before conception, at conception, during each trimester of pregnancy, and after delivery).

Panel 2A. Linear representation of the complement levels overtime in the entire cohort of systemic lupus erythematosus patients. Panel 2B. Linear representation of the fluctuations of complement levels during pregnancy in patients with systemic lupus erythematosus with and without the occurrence of flares during pregnancy. Panel 2C. Linear representation of complement levels during time in patients with and without lupus nephritis (LN). Panel 2D. Linear representation of complement levels during time in patients with and without LN and presence, or absence, of flare during pregnancy.

Table 1. Complement levels at the six different time-pcints (values expressed as mean ±SD), according to diagnosis of lupus nephritis (LN) or presence of a disease flare during pregnancy.

Table 2. Odds Ratios according to diagnosis or preserior of flare and different ΔC3 levels.

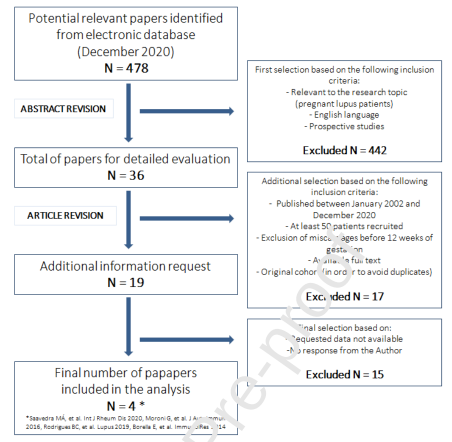


Figure 1. Literature search strategy

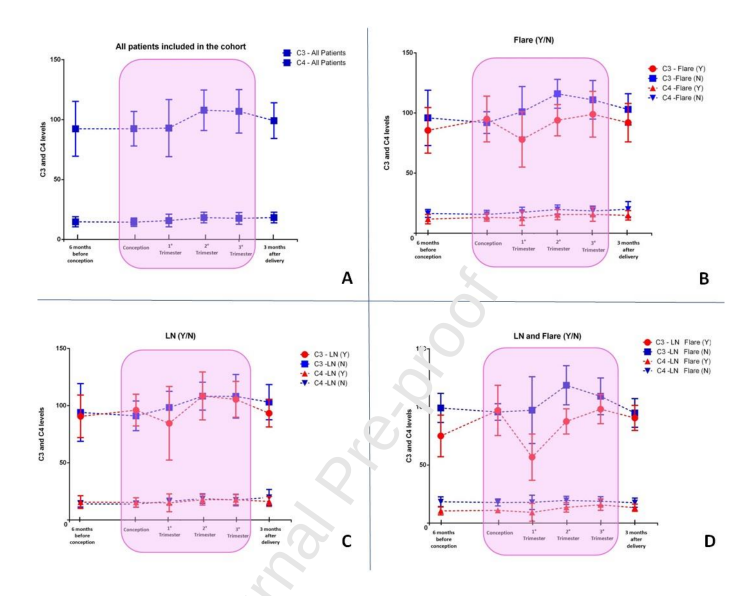


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	All SLE	Patients	Patients	Patients with	Patients without	Patients with LN	Patients with LN and
	patients	with LN	without LN	flare	flare (N=362)	and flare (N=73)	without flare
	(N=532)	(N=237)	(N=295)	(N=170)			(N=164)
C3 6 months							
before	92.3±22.9	90.7±18.6	94.1±25.2	85.6±19.1	95.6±23.3	75 ±17.9	99.1±12.5
pregnancy							
C3 at conception	92.4±14.4	96.1±13.9	91.1±13	95.3±19.5	91.8±9.1	97 ±21.6	95.6±7.1
C3 T1	92.9±23.8	84.6±32.2	98.4±14.1	78.3±22.8	100.5±20.7	56.8 ±19.9	97.2±28.7
C3 T2	107.8±16.9	108.5±21	108.3±12.2	94.16±13.4	115.7±12.3	87.5 ±10.9	118.6±16.8
C3 T3	106.9±18.1	105.5±15.7	108.2±19.1	98.97±18.6	111.4±16	98.1 ±12.6	109.1±15.8
C3 3 months PP	99.1±14.9	93.4±12	103.1±15.4	92.4±15.7	102.6±13.4	90.5 ±10.8	94.8±12.3
C4 6 months							
before	14.7±4.2	15.7±5.5	14.1±2.8	11.8±3.9	16. <mark>! ±3.3</mark>	10.5±3.4	18.4±4.2
pregnancy							
C4 at conception	14.4±3.5	15.4±4.1	13.9±2.8	13.3±3.2	15 7±3.4	11±1.3	17.8±3
C4 T1	15.8±5.3	15±7.8	16.3±2.8	12.5±5.9	1 5±4.2	9.3±7.6	17.9±6.2
C4 T2	18.3±4.4	17.7±4.7	18.7±4.2	15.5±4.?	19.8±3.7	13.6±4.1	19.6±3.5
C4 T3	17.6±4.9	17.8±4.4	17.5±5.1	15.7:5.8	18.6±4	15.8±4.8	18.8±3.9
C4 3 months PP	18.3±6.2	16.2±4.3	19.8±6.9	14.° ±3.9	20±6.4	13.3±3.1	17.6±4
ΔC3(ΔC3 T1—at conception)	10.3±43.2	0.5±53.2	16.6±34.3	-6.7.48.8	18.8±37.6	-36.1±42.6	17.3±49.1
ΔC4 (ΔC4 T1– at conception)	3.4±7.6	1.5±9.1	4.5±6.3	1.2±8.1	4.4±7.1	-1.1±8.5	2.8±9.1

Table 1. Complement levels at the six different timepoints (values expressed as mean ±SD), according to diagnosis of lupucine phritis (LN) or presence of a disease flare during pregnancy

Results highlighted in boluare statistically significant.

Abbreviations: SLE, systemic lupus erythematosus; LN, lupus nephritis; T1, 1^{st} trimester of gestation; T2, 2^{nd} trimester of gestation, T3, 3^{rd} trimester of gestation; PP, post-partum period (up to 3 months after delivery).

	FLARE	FLARE	LN	LN	LN & FLARE	LN & FLARE
	OR	CI 95%	OR	CI 95%	OR	CI 95%
ΔC3 ≥15 mg/dL	0.3	0.2-0.5	1.2	0.7-2.5	0.06	0.02-0.3
ΔC3 ≥10 mg/dL	0.5	0.3-0.7	0.4	0.3-0.5	0.03	0.01-0.1
ΔC3≥5 mg/dL	0.4	0.3-0.6	0.2	0.1-0.3	0.02	0.01-0.07
ΔC3 no change defined as	1.1	0.6-1.9	2.2	1.3-3.6	5.2	2.9-9.3
[-2;+4] mg/dL	1.1	0.6-1.9	2.2	1.3-3.0	5.2	2.9-9.5
ΔC3 ≤5 mg/dL	3.1	2.1 -4.8	6.1	3.9-9.6	6.5	3.9-11.2
ΔC3 ≤10mg/dL	3.3	2.2-5.1	7.2	4.5-11.7	5.6	3.3-9.7
ΔC3 ≤15mg/dL	3.9	2.5-6	6.4	4-10.3	6.2	3.6-10.7

Table 2. Odds Ratios (OR) according to lupus nephritis (LN) diagnosis or presence of flare and different ΔC3 levels (first trimester –at conception)

Results highlighted in bold are statistically significant.

Declaration of interests

☑The authors declare that they have no known competing financial interests or personal relationships that
could have appeared to influence the work reported in this paper.

☐ The authors declare the following financial interests/personal relationships which may be considered as potential competing interests:

Highlights

C3/C4 values progressively increase in lupus pregnancy as in physiological pregnancy, but the peak is in the second trimester, then there is a decline, particularly in C3.

- Patients with flare during pregnancy and those with previous LN showed significantly lower complement levels than those who did not have flare and did not have renal involvement, respectively.
- The lowest complement levels pre-pregnancy and at conception were observed in patients with previous LN and related gestational flare.
- The fluctuations of C3 and C4 levels in the first trimester versus conception displayed the highest clinical significance in predicting both SLE flares and APO as compared to variations between other timepoints.
- A decrease, stable values, or an increase smaller than 5 mg/dL in C3 levels between conception and the first trimester were all factors associated with an increase in the risk of SLE flares and APO.
- Validation studies will clarify these findings and define the ole of C3 and C4 levels during the preconception period/early pregnancy as tools for prerioting SLE flares and APO and implementing individualized treatment strategies.
- In lupus patients, C3 and C4 monitoring should be incorporated into the clinical assessment before conception and throughout the entire pregnancy.