Title: HYDROXICLOROQUINE BLOOD CONCENTRATION IN LUPUS NEPHRITIS: A DETERMINANT OF DISEASE OUTCOME?

OR:

ASSOCIATIONS BETWEEN BLOOD LEVELS OF HYDROXYCHLOROQUINE AND OUTCOME IN LUPUS NEPHRITIS

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ABSTRACT

Introduction: Hydroxychloroquine (HCQ) is a recommended drug in Systemic Lupus Erythematosus (SLE). It has a long terminal half-life, making it an attractive target for therapeutic drug monitoring. The aim of this study was to establish a relationship between blood HCQ concentration and lupus nephritis activity.

Methods: Retrospective observational study with data collected from clinical and laboratory records. Inclusion: Patients followed in Lupus Clinic with biopsy proven ISN/RPS class III, IV or V lupus nephritis, on HCQ for at least 3 months (200-400mg daily) & with HCQ levels measured during treatment. Excluded: Patients on renal replacement therapy at baseline or lost to follow-up.

Results: In 171 patients, HCQ level was measured in 1282 samples. Mean (+/-SD) HCQ blood level was 0.75(+/-0.54)mg/L and it was bimodally distributed. HCQ level less than 0.20mg/L (232 samples, 18.1%) appeared to define a distinct group of abnormally low HCQ levels. For patients in complete or partial remission at baseline, compared to those remaining in remission, patients with renal flare during follow-up had significantly lower average HCQ level (0.59 vs 0.81mg/L, p=0.005). Our data suggests an HCQ target level to reduce the likelihood of renal flares greater than 0.6mg/L [600ng/mL] in those patients with lupus nephritis.

Conclusion: HCQ level monitoring may offer a new approach to identify non-adherent patients and support them appropriately. We propose a HCQ minimum target level of at least0.6mg/L to reduce renal flare rate, but it will require a prospective study for validation.

KEYWORDS: Lupus Nephritis, Disease Activity, Treatment, Glomerulonephritis, Systemic lupus erythematosus
INTRODUCTION

Lupus nephritis is a major complication of systemic lupus erythematosus (SLE), occurring in up to two thirds of SLE patients. Renal remission is important, with those patients that enter complete remission having a higher renal, and overall survival compared to those patients in whom remission is not achieved.[1] Even a partial remission in lupus nephritis is associated with a significantly better patient and renal survival compared with no remission.[2-5] Renal flares average about 8 per 100 patient years for the first five years after attaining remission.[6] Severe disease at baseline, a delay in reaching remission, and attainment of partial compared to complete remission are some of the factors predisposing to renal flares.[6-8] Non adherence to lupus nephritis treatment is the most important factor for non-remission or renal relapses.[9-11]

Hydroxychloroquine (HCQ) is currently recommended long-term for all patients with SLE provided no contraindications exist. There is evidence of multiple beneficial effects, including control of disease activity by its immunomodulatory effects, reduction of cardiovascular events, lipids and glucose lowering effects, antithrombotic effects and improved survival.[12-16]

HCQ has a long terminal half-life of over 40 days and a steady-state level in blood with small daily fluctuation,[17] making it an attractive target for therapeutic drug monitoring to evaluate adherence to medication as well as efficacy.

A pharmacokinetic/pharmacodynamic relation for blood HCQ concentration has been found in rheumatoid arthritis and cutaneous lupus.[18-20] For SLE, low whole-blood HCQ concentrations suggest non-adherence and may predict disease exacerbations, with one study showing that a cutoff of 1mg/L had a negative predictive value of 96% for a systemic flare during follow-up.[21] There is also evidence of a protective effect of hydroxychloroquine in retarding renal damage occurrence in SLE.[22] but there are no reports to our knowledge looking specifically at HCQ blood levels in patients with lupus nephritis.

The aim of this study was to establish a relationship between blood HCQ concentration and lupus nephritis activity, including its relation with time to remission and risk of renal flares.
METHODS:

Study design: This is a retrospective observational study with data collected from clinical and laboratory records from January 2011 to October 2015. This study was designed, implemented, and reported in accordance with ICH guidelines for Good Practice and Declaration of Helsinki 1975, revised Hong Kong 1989. Since the standard clinical protocol was followed for patient management, it was decided in discussion with Chair of the local Research Ethics Committee that there was no requirement for formal ethical review. The manuscript was prepared in accordance with the STROBE guidelines for observational studies.

Setting: Patients were recruited from those attending the Imperial College Healthcare NHSTrust Lupus Centre outpatient and inpatients services, Hammersmith hospital, London, UK. Patients had follow up appointments according to clinical indications. We currently follow up approximately 400 patients with lupus (the majority of whom have had at least one episode of lupus nephritis), not all of whom are on HCQ.

Inclusion criteria: All patients with age ≥18 years with biopsy proven ISN/RPS class III, IV or V lupus nephritis, on HCQ for at least 3 months and with HCQ levels measured during treatment.

Exclusion criteria: Patients on renal replacement therapy at baseline or lost to follow-up.

Outcome variables: The baseline of the study was defined as the date of first measurement of HCQ concentration in blood. The definitions for complete remission, partial remission and renal relapse were taken from the study by Condon M et al previously performed in this centre.[23]Complete remission was defined by the combination of urine protein/creatinine ratio (uPCR) of less than 50 mg protein/mmol creatinine and serum creatinine no greater than 15% above baseline. Partial remission was defined by the combination of uPCR<300 mg/mmol with >50% reduction from baseline and serum creatinine not more than 15% from baseline. Renal relapse was defined as a persistent increase of >30% in proteinuria and/or serum creatinine, requiring renal biopsy or increase/change in immunosuppression. Outcome variables were assessed at baseline and at each follow-up visits.

Exposure variables: All patients were on HCQ and the standard clinical practice was to administer 400 mg in one or two divided doses. The formulation was not designated and many patients take generic HCQ. The dose of the drug was decreased by the treating physician if drug toxicities occurred. HCQ blood level was tested at least three months after starting the drug. Ophthalmological testing was undertaken on a case to case basis and drug toxicities were recorded as and when reported. Concomitant immunosuppression was continued as per local standard of care regimens.
Predictor variables: Demographic, clinical, laboratory and pharmacokinetic data were collected at baseline and at follow-up visits. Demographic variables were age, gender and race. Clinical variables were weight, ISN/RPS class of lupus nephritis, co-morbidities, duration and time of diagnosis of SLE and lupus nephritis and use of immunosuppression. Laboratory variables included uPCR, serum creatinine, serum albumin, haemoglobin, total leukocyte count, C reactive protein and SLE serology (levels of anti dsDNA antibody, C3 and C4). Pharmocokinetic variables were HCQ dosing, HCQ blood levels and concomitant mycophenolic acid (MPA) blood levels if available.

Data sources: Demographic, clinical and HCQ pharmacokinetic details were collected from patient records, the hospital database and from Renal Clinical Leslie Brent Laboratory. Data were collected for all patients for all follow-up visits during the study period.

Whole blood HCQ concentrations were determined by an adaptation of our previously published liquid chromatography - tandem mass-spectrometric method using HCQ-d₃ as the internal standard.[24] Total imprecision was 8.3% @ 0.25mg/L, 4.2% @ 0.90mg/L and 4.1% @ 2.1mg/L (n=72). The measurement of whole blood rather than plasma hydroxychloroquine is important. Whole blood concentrations are approximately five times the plasma concentrations, are more precise, and are favored for pharmacokinetic measurements.

Study sample size: The number of consecutive cases that met the inclusion criteria within the study period (January 2011 to October 2015) determined the sample size.

Statistical methods: Individual HCQ levels are presented as histograms for inspection. Undetectable HCQ levels (< 0.1mg/L) were recorded as 0.05mg/L for analysis. Within patient variability in HCQ level was assessed by regression model with patient as an independent predictor variable. Average HCQ level for each patient was used for outcome analysis, with student t test and chi square test as appropriate for numerical and categorical comparisons.
RESULTS:

Participants: 171 patients > 18 years of age with ISN/ RPS class III, IV and/or V, on HCQ for at least 3 months and had HCQ blood levels measured during treatment. The mean follow-up period was 26.9 +/- 15.7 months.

Descriptive data: The baseline demographics, clinical and laboratory parameters are summarized in Table 1. The majority of patients were women (n=147, 86%) and the mean age was 39.8±15.6 years. A total of 73 patients (42.7%) were of South East Asian origin from the Indian subcontinent.

At baseline out of 171 patients, 91 (53.2%) were in complete remission from their prior lupus nephritis, 36 (21.1%) were in partial remission and 39 (22.8%) had active lupus nephritis.

Outcome data: HCQ level was measured in 1282 samples from the 171 patients and mean (+/-SD) HCQ blood level was 0.75(+/-0.54)mg/L.

We have looked at patient factors affecting HCQ levels and found lower HCQ concentrations in heavier patients (R=-0.179, p=0.019) (Figure 1), but no relation to age, gender or ethnicity (data not shown).

Levels whilst taking 200mg/day were lower than whilst taking 400mg/day (0.58+/-0.47 vs 0.79+/-0.54mg/l, p<0.001). Over the study period, 74.9% of patients took 400mg/day throughout, 14.0% took 200mg/day or less, with the remaining 11.1% taking 200 or 400mg/day during different periods. There was no defined protocol for dose selection, but those on less than 400mg/day for part of the study were on lower baseline doses of prednisolone (2.2 vs 3.7mg/day, p=0.020) and mycophenolate mofetil (0.62 vs 1.01g/day, p=0.005) and had lower eGFR (68 vs 78, p=0.020), but they were not different in terms of age, gender, ethnicity, body weight or serum albumin.

HCQ levels were bimodally distributed: HCQ level less than 0.20mg/L (232 samples, 18.1%) appeared to define a distinct group of abnormally low HCQ levels, which suggests a definition of non-adherence to the treatment though in some cases it might be explained by insufficient dosing as an effect of dose was noted (Figure 2).

Less than 40% of total variation in HCQ levels occurred within individuals, suggesting consistency of HCQ level over time (Figure 3). In those patients (n=26, 15.2%) with average HCQ level less than 0.2mg/L (poorly adherent patients), 88.7% of individual samples were also less than 0.2mg/L. Average HCQ level for each patient was therefore used for clinical correlations.

Regarding clinical outcome, for those with active nephritis at baseline, average HCQ levels in those who subsequently achieved remission were similar to those who remained active (p=0.23). In
contrast, for patients in complete or partial remission at baseline, compared to those remaining in remission, patients with renal flare during follow-up had significantly lower average HCQ level (0.59 vs 0.81mg/L, p=0.005). (Figure 4)

Considering only the patients in complete or partial remission at baseline, the proportion with disease flare according to quintile of average HCQ level is shown in Figure 5(Q1: <0.31mg/L, Q2: 0.31-0.62mg/L, Q3: 0.63-0.81mg/L, Q4: 0.81-1.12mg/L, Q5: >1.12mg/L). The p value (chi square test) is 0.2, but a significant difference is seen when comparing Q1-2 with Q3-5 (p = 0.041). The Q2 / Q3 boundary is 0.62 mg/L, which can be considered the lower end of the desirable HCQ blood target level.

HCQ level over 0.6mg/l was achieved in 68.8% of patients taking 400mg/day throughout, and 46.5% of patients on lower doses for part of the time. A starting dose of 400mg/day therefore seems reasonable though reductions or increases may be appropriate for a significant number.

HCQ levels were left-skewed, with the highest quintile ranging from 1.1 to 3.6mg/L. Toxicity had not been systematically recorded so cannot be quantified: the most common problem noted was skin pigmentation but a prospective study will be required to relate this to drug levels. No major toxicity was recorded.

Patients in the top quintile of average HCQ level experienced fewer flares during follow up, and no flattening of the level-outcome slope was observed. It is therefore not possible to suggest an upper limit for a target range based on this study. When faced with high levels and possible toxicity however, clinicians may find it helpful to know that average HCQ level over 1.6, 1.8 and 2.0mg/l was seen in 4.7, 1.8 and 0.6% of patients (8, 3 and 1 patient). The HCQ benefits observed in this study were therefore achieved with levels lower than these in the majority of patients.

Our data therefore suggest an HCQ target level to prevent flares should be at least 0.6mg/L [600ng/mL].
DISCUSSION

HCQ is recommended long-term for all patients with SLE provided no contraindications exist.[12-16] Its mechanisms of action are still under investigation, but there is evidence that HCQ, by blocking the toll-like receptor 7 and 9 in plasmacytoid dendritic cells, inhibits interferon-alpha production, which plays a crucial role in SLE pathogenesis.[25] It has a long terminal half-life of over 40 days with small daily fluctuation of the blood levels, making it an attractive target for therapeutic drug monitoring to evaluate adherence to medication as well as efficacy.

The blood HCQ concentration varies widely among patients with SLE. However, current information on factors influencing this variability is still controversial. A retrospective analysis of data from the Plaquenil Lupus Systemic (PLUS) study found that high body mass index, no treatment with corticosteroids, increased delay between the last tablet intake and measurement of blood HCQ concentration, low platelets and neutrophil counts, and high estimated creatinine clearance were associated with low blood HCQ concentration.[26] The authors did not find an association of ethnicity or smoking and blood HCQ concentration and no pharmacokinetic drug-drug interaction with antacids or inhibitors and inducers of cytochrome P450 enzymes.[26] In contrast, another study did not find a relation between HCQ levels and eGFR, but only 14% of patients had stage 4 or 5 CKD.[27] This and other studies also did not find any significant relationship between HCQ levels and gender, body weight, body mass index, renal function or chronic smoking.[27-29] On the other hand, a recent Asian study reported that HCQ levels were related to CYP2D6 polymorphisms in Korean lupus patients taking oral HCQ, which may explain why there is wide variation in blood HCQ concentrations.[30] In our study, HCQ levels were lower in heavier patients but unaffected by age, gender or ethnicity. It was also interesting to see that some patients on lower daily doses of HCQ (200 mg daily) achieved adequate serum levels of HCQ, as shown in figure 2, highlighting the variability between patients. So, for some patients 400 mg daily would be a higher dose than necessary.

Poor adherence to therapeutic regimens is a common problem in patients with chronic diseases including SLE and is associated with a higher risk of flares, morbidity, hospitalisations and poor renal outcome. The rates of non-adherence in SLE patients range from 3% to 76% depending on the assessment methods, which are all subject to limitations.[31] Costedoat-Chalumeau N et al reported in 2007 a total non-adherence with HCQ of 7% (undetectable levels). Reasons for non-adherence included both those attributed to HCQ characteristics (concern about potential side effects, perceived inefficacy of the drug compared to other treatments, adverse side effects attributed to the drug) and related to patients characteristics (failure to accept the disease and forgetfulness).[32] In a cohort of 70 patients with childhood-onset SLE, only 32% of patients were sufficiently adherent to HCQ.[33] A cross-sectional study described that 13% of SLE patients had whole blood HCQ levels of < 15 ng/mL (complete non-adherence).[28] A more recent study found a proportion of total non-adherence of 11%,
defined as HCQ serum level of < 10 ng/mL.[27] Our study showed that an HCQ level lower than 0.2mg/L [200ng/mL] can be a useful indicator of poor adherence, which may reflect a more generalised adherence problem involving other disease modifying agents. Using this definition, we found non-adherence with HCQ consistently occurs in at least 15.6% of our lupus nephritis population. However, we cannot exclude that in some cases these low levels were a result of under-dosing or differences in HCQ pharmacokinetics, instead of non-adherence. Direct comparison between studies is confounded by different cut-off values used to define non-adherence and differences in the HCQ assay. For patients identified as being poorly adherent by therapeutic monitoring, counselling may improve medication adherence. In the study by Costedoat-Chalumeau N et al, adherence improved after patients were informed of the results and interviewed about their adherence to treatment in a non-judgmental discussion.[32] Additionally, there is evidence from a cross-sectional study that counseling patients with low HCQ levels led to an increase in its concentration, where only 56% of the patients had levels above 500 ng/ml [0.5mg/L] at their first follow-up visit versus 80% at last follow-up in those who attended 3 visits or more.[28]

We collected mycophenolic acid (MPA) levels on a number of the patients included in this study. However, whereas HCQ levels are little affected by time of last dose, this is not the case for MPA, and few levels were true troughs. So, from a retrospective data collection, where timing of dose was not routinely collected, it is impossible to draw any firm conclusions as to whether there is generalised or specific non-adherence. Some patients appear to take neither their MMF or HCQ (repeatedly undetectable levels of both) whereas some definitely take one and not the other. We have reviewed the baseline MPA and HCQ levels – 102 patients had simultaneous MPA and HCQ levels. Of those with HCQ levels <0.2 (n=17) just one (5.9%) had an undetectable level of MPA, 9 (52.9%) had levels <1.0 (which could well represent a long trough) and 10 (58.8%) had levels below the minimum therapeutic level (1.4). In the 85 patients with HCQ levels >or=0.2, none had undetectable MPA levels, 38 (44.7%) had levels <1.0 and 57 (67.1%) had levels below 1.4. Conversely, 41.2% of those with undetectable HCQ had therapeutic levels of MPA, vs just 32.9% of those with HCQ levels >0.2. This suggests that non-adherence is not necessarily uniform.

The benefits of HCQ treatment in SLE have been demonstrated in a randomized, double-blind, placebo-controlled study of 47 SLE patients, in which the risk of clinical SLE flares rose 2.5-fold during a 6-month period after discontinuation of the drug.[34] There is evidence that whole-blood HCQ concentrations suggest non-adherence and may predict disease exacerbations, with one study showing that a cutoff of 1000ng/mL [1mg/L] had a negative predictive value of 96% for flare during follow-up.[21] This cutoff was in the same range as those able to block intracellular TLRs in vitro, currently considered the key activity of HCQ in SLE.[25] In contrast to these results, two other studies did not find a significant concentration-effect relationship for HCQ in the treatment of
However, in the first study consistently high enough blood HCQ concentrations were not achieved.[35] In contrast, in the second study, in a subgroup of patients with serological and clinical remission and having therapeutic HCQ levels (defined as > 500 ng/mL), a trend of lower disease activity and incidence of flares was indeed observed.[27] In 2013, Costedoat N et al showed that low HCQ concentration is associated with higher SLE activity, but adjusting the HCQ dose to a target level in blood did not reduce flares over a 7-month follow-up.[36] However, the maintenance of [HCQ] above 1000 ng/ml [1mg/L] during the 7-month follow-up was difficult to achieve.[36] 

There is evidence of a protective effect of hydroxychloroquine in retarding renal damage occurrence in SLE and protecting against renal insufficiency.[22] Additionally, HCQ was previously shown to be an independent predictor of complete renal remission in SLE patients treated with mycophenolate mofetil for membranous lupus nephritis.[37] However, there are no reports to our knowledge looking specifically at HCQ blood levels in patients with lupus nephritis as in our study. We have shown that for patients with active nephritis at baseline, average HCQ levels in those who subsequently achieved remission were similar to those who remained active. In contrast, for patients in complete or partial remission at baseline, compared to those remaining in remission, patients with renal flare during follow-up had significantly lower average HCQ levels (0.59 for patients who flared vs 0.81mg/L for patients who maintained remission). Therapeutic drug monitoring may therefore assist in maintaining remission in lupus nephritis. Of course it is possible that remission was maintained by adherence to other maintenance immunosuppression in those who also took their HCQ. But this does not diminish its value in evaluating adherence, whether specific or more generalised.

Our retrospective data also provide a suggestion for a therapeutic target range of HCQ levels. Renal flares were significantly fewer in those with HCQ levels > 0.6 mg/L, which can be considered the lower end of the desirable HCQ target level. It is not possible from our study to define an ideal upper level as toxicity was not systematically monitored, a limitation of our study.

In general, HCQ is well tolerated and rarely needs to be discontinued for an adverse reaction.[16] Gastrointestinal intolerance, pruritus and other cutaneous manifestations are not rare, but usually disappear with dose reduction. Potential severe side effects include retinal, neuromuscular and cardiac manifestations. Although these are rare, they require immediate discontinuation of treatment. In our study, no significant adverse effects were found even for high HCQ levels, but further studies are necessary to confirm whether the high levels seen in some are necessary. The most common problem in our study noted was skin pigmentation but a prospective study will be required to relate this to drug levels.

Another limitation of this study includes the fact that HCQ samples were collected when patients attended clinic, so intervals between levels were not consistent between patients. That may result in
ascertainment bias as patients with poor outcomes are likely over-represented in the analysis of the cohort as a whole. However, all patients with available HCQ blood levels were included even if only one measurement was available, and there was significant consistency in levels over time for patients tested more than once, minimizing its effect.

Finally, the retrospective and observational nature of this study is the main limitation to our conclusions. A prospective study is necessary to validate our results.

In conclusion, non-adherence remains a major cause of poor outcomes in patients with lupus nephritis, and our study suggests it consistently occurs in at least 15.6% of our lupus nephritis population, at least with respect to HCQ. HCQ level monitoring offers a new approach to identify such patients and support them more appropriately. We propose a minimum HCQ target level to prevent flares of 0.6mg/L [600ng/mL], but it will require a prospective study for validation.
Contributions: LL conceived the idea; GC and JL developed and validated the assay. CC and SA contributed equally to collecting data and drafting the manuscript. All authors contributed to the study design, data analysis and manuscript preparation.

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There are no competing interests.

Ethical approval information: Since the standard clinical protocol was followed for patient management, it was decided in discussion with Chair of the local Research Ethics Committee that therefore there was no requirement for formal ethical review.

I confirm this manuscript has not been accepted for publication elsewhere, is not being considered for publication elsewhere and does not duplicate material already published. I confirm all authors consent to publication of this manuscript.
REFERENCES


### TABLES

#### Table 1

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*Current treatment at time of baseline measurement of HCQ.*
Figure 1 - Variation of hydroxychloroquine levels according to patient weight (HCQ –hydroxychloroquine)

\[ y = -0.0055x + 1.1281 \]
\[ R^2 = 0.03166 \]

Figure 2 - Distribution of total hydroxychloroquine (HCQ) blood levels measured. Black - Daily HCQ dosage of 200 mg. Grey - Daily HCQ dosage of 400 mg.
Figure 3 - Consistency of hydroxychloroquine (HCQ) blood levels over time per patient.

Figure 4 - Average hydroxychloroquine (HCQ) blood level by follow-up status (complete remission vs flare during follow-up in those in complete or partial remission at baseline).
Figure 5 - Percentage of lupus nephritis flare during follow-up for patients in complete/ partial remission at baseline per average hydroxychloroquine blood levels quintile Q1: <0.31mg/L, Q2: 0.31 - 0.62mg/L, Q3: 0.63 - 0.81mg/L, Q4: 0.81 - 1.12mg/L, Q5: >1.12mg/L.
Conflict of Interest Statement: none declared

[Signature]

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