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eprints@whiterose.ac.uk https://eprints.whiterose.ac.uk/ 1 Hydroxychloroquine effectiveness in reducing symptoms of hand osteoarthritis: a

2 Randomized Trial

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- 45 trial
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49	Abstract
50	Background: It is thought that synovitis may play a role in producing symptoms in people
51	with hand osteoarthritis (OA), but data on slow-acting anti-inflammatory treatments are
52	sparse.
53	
54	Objective: To determine the effectiveness of hydroxychloroquine versus placebo as an
55	analgesic treatment for hand OA.
56	
57	Design: Randomized, double-blind, placebo-controlled clinical trial with 12-month follow-up.
58	
59	Setting: 13 primary- and secondary-care centres in England.
60	
61	Participants: Of 316 patients screened, 248 participants (82% women, mean age 62.7
62	years) with symptomatic (VAS pain ≥4/10) and radiographic hand OA were randomized. 210
63	(84.7%) completed the 6-month primary endpoint.
64	
65	Intervention: Hydroxychloroquine (200-400mg) or placebo (1:1) for 12 months in addition to
66	ongoing usual care.
67	
68	Measurements: The primary endpoint was average hand pain during the previous 2 weeks
69	(numerical rating scale [0-10], NRS) at 6-months. Secondary endpoints included self-
70	reported pain and function, grip strength, quality-of-life, radiographic structural change and
71	adverse events. Baseline ultrasonography was performed.
72	
73	Results: At 6 months, the mean hand pain (as measured by NRS) was 5.49 and 5.66 in the
74	placebo and hydroxychloroquine groups, with a treatment difference of -0.16 points (95% CI:
75	-0.73 to 0.40, p=0.57). Results were robust to adjustments for adherence, missing data and
76	use of rescue medication. There were no significant treatment differences at 3, 6 or 12-

77	months for any secondary outcomes. On ultrasound, 94% (133/143) had \geq 1 joint positive for
78	greyscale synovitis, 59% were Power Doppler positive. Baseline structural damage or
79	synovitis did not affect treatment response. Fifteen serious adverse events were reported
80	(hydroxychloroquine: 7 [3 defined as possibly related], placebo: 8).
81	
82	Limitations: Hydroxychloroquine dosage restrictions may have reduced efficacy.
83	
84	Conclusions: Hydroxychloroquine was no more effective than placebo for pain relief in
85	people with moderate to severe hand pain and radiographic OA.
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87	
88	
89	Trial Registration: ISRCTN91859104
90	Funding Source: Arthritis Research UK Clinical Studies Grant (19545)
91	
92	

93 Symptomatic hand osteoarthritis (OA) affects 4-31% of adults over the age of 70, and 3-15% over the age of 60 (1-7). Individuals report chronic persistent pain and considerable difficulty 94 with daily activities (8). However there are few effective therapies for this condition and use 95 of these therapies is often limited by patients' comorbidities or toxicities (9-11). Consequently 96 97 primary and secondary care physicians seek alternative options to improve quality of life for people with this painful, disabling disease. Anecdotal reports suggest hydroxychloroguine 98 (HCQ) is one such therapy. It has been used as an unlicensed treatment in many countries 99 100 when other options have failed, mainly for the subset of patients with "inflammatory" hand OA (12,13). HCQ is an established drug treatment for inflammatory arthritides such as 101 102 rheumatoid arthritis (RA), supported by placebo-controlled trials demonstrating its efficacy, as a monotherapy and in combination with other RA drugs, and acceptable safety profile 103 104 (14,15). With increasing evidence that inflammation is highly prevalent in OA and may have 105 a role in symptoms (16-20) and three small pilot studies suggesting reduction in hand pain 106 with HCQ (21-23), there is a rationale for exploring the efficacy of HCQ as a treatment for 107 hand OA.

108

The objective of the Hydroxychloroquine Effectiveness in Reducing symptoms of hand
Osteoarthritis (HERO) Trial was to test the hypothesis that HCQ is an effective symptomatic
treatment when used in people with at least moderate symptomatic hand OA and inadequate
response to current therapies including NSAIDs and opioids.

113 Methods

114 **Design Overview**

115 The HERO trial was an investigator-led, pragmatic, multi-centre, superiority, randomized, 1:1 116 placebo-controlled trial. The research protocol (Appendix 1) was approved by Leeds East 117 Research Ethics Committee (12/YH/0151), the UK Medicines and Health Regulatory 118 Authority (MHRA) and registered on ISRCTN (ISRCTN91859104) in parallel. Participants were recruited from September 24th 2012 until May 27th 2014, with participants followed-up 119 120 for 12-months post-randomization (follow-up completed April 25th 2015). Written informed 121 consent was obtained for all participants prior to screening. One participant was recruited (24.09.2012) prior to protocol registration (17.10.2012), however no changes were made to 122 the protocol between these time-points and therefore this participant is similar to all other 123 trial participants. Full trial design details are available (Appendices 1-4). 124

125

126 Setting and Participants

The trial involved 13 National Health Service (NHS) hospitals in England, with recruitment 127 taking place through primary care and secondary care-based musculoskeletal clinics. 128 129 Patients were eligible if aged ≥18 with self-reported, inadequate response or side-effects to existing medication (including paracetamol, oral NSAID or opioid); moderately severe 130 symptoms (hand pain ≥4/10 on a 0-10 visual analogue scale) for more than half of days in 131 the last 3 months; fulfilled American College of Rheumatology criteria for OA (24); hand 132 radiographs in the past 5 years with changes consistent with OA; stable, no change to or no 133 134 use of analgesics (including NSAIDs) for at least 4 weeks or glucosamine or chondroitin for 135 at least 4 months; and capable and willing to give consent and adhere to the study protocol. 136 Exclusion criteria were inflammatory arthritis; psoriasis; CMC joint (CMCJ) involvement only 137 or predominant CMCJ pain; oral, intramuscular, intra-articular, intravenous steroids or other anti-synovial agents or any new hand OA therapies during the last two months; intra-articular 138 hyaluronans in last 6 months; uncontrolled disease states where flares are commonly 139 140 treated with corticosteroids; serious uncontrolled medical condition; unexplained visual

impairment; pregnant or lactating; melanoma or non-skin cancer in the past 3 years,

significant haematological or biochemical abnormality (Appendix 4). Rheumatoid factor (RF)

- and anti-CCP were measured in all eligible participants to exclude inflammatory arthritis.
- 144

145 Randomization and Interventions

146 Patients were randomized to either hydroxychloroquine (200, 300 or 400mg, with dosage 147 calculated according to ideal body weight to give a maximum dose of 6.5mg/kg/day) or 148 placebo. Randomization (1:1) was computer-generated (PRISYM ClinTrial) in advance by 149 the contract manufacturer using random permuted blocks, without stratification. The contract manufacturer prepared trial drug with over-encapsulation to create identical intervention and 150 placebo-control products with no involvement from the research team, and assigned 151 intervention and control drug packs in sequence to recruiting sites. All parties remained blind 152 153 to treatment allocation throughout the trial. Adverse events, vital signs and blood monitoring were assessed on an ongoing basis during follow-up. All elements of participant care were 154 left to the discretion of the site research team in line with the pragmatic nature of the HERO 155 trial, with the exception that steroids and new or experimental interventions were not 156 157 permitted during follow-up. Adherence to trial medication was collected using multiple methods to provide an estimate of compliance, including site-reported non-adherence, 158 participant-reported Brief Medication Questionnaire (25), and pharmacy records of returned 159 medication. Quality of adherence data was reviewed prior to unblinding to determine non-160 adherence criteria for analysis (Appendix 4). Participants were asked about adverse events 161 (AEs) at all visits and these were reviewed by a physician for severity, duration and 162 relatedness to investigational medicinal product (IMP). SAEs were defined according to pre-163 164 specified criteria, as detailed in the protocol (Appendix 1), assessed for causality and 165 expectedness by a physician and reported within 24 hours.

166

167 Outcomes and Follow-up

168 Data collection was completed using standardized case report forms at screening, baseline,

169 3, 6 and 12-months. The primary outcome was overall hand pain severity over the past 2 weeks, measured on an 11-point (0-10) Numerical Rating Scale (NRS), at 6-months follow-170 up (26). This outcome was also assessed at baseline, 3 and 12-months. Secondary 171 outcomes included: pain severity in the most painful joint (NRS over last 2 weeks), AUSCAN 172 173 pain and function scales (27), grip strength (measured using a dynamometer) (28), structural 174 damage using bilateral hand radiograph data (29), Osteoarthritis Quality of Life (OAQoL) (30), and Short-form 12 (SF-12) Physical and Mental Component Score (31). Bilateral hand 175 176 radiographs (baseline, 12-months) were captured according to a standardized protocol 177 (Appendix 4) and scored in pairs at the end of the study by a musculoskeletal radiologist who was blinded to participant identity and treatment allocation. Baseline ultrasound imaging 178 was performed for the dominant hand of all participants enrolled at the six ultrasound sub-179 study centres using a standardised protocol (Appendix 4) and following a group training day 180 181 for the ultrasound operators.

182

A full list of secondary outcomes is described in Appendix 4 and Appendix Table 1. Costeffectiveness data, collected at baseline and 12-months, will be presented in a separate publication.

186

187 Statistical Analysis

The HERO trial was powered to detect a standard effect size of 0.4, equivalent to the reported effect size of NSAIDs as a treatment for hand OA (32,33) and a reduction in pain of 0.8 score points (or 15%) on the NRS (32,33) which lies within the minimal clinically important difference for change in pain in a randomized trial (10/20%)(34). To detect a standard effect size of 0.4 with 80% power and 5% two-sided significance, 99 patients were required per arm. Allowing for 20% dropout and equal numbers per centre, the total target sample size was 252 patients.

195

196 The analyses followed a pre-specified statistical analysis plan, endorsed by the data and safety monitoring committee, and were performed using Stata version 13 (StataCorp, Texas, 197 198 USA). The statistician remained blinded to treatment allocation until verification of the 199 primary analysis. The primary analysis was intention-to-treat (ITT), analysing participants in 200 their randomization group. A linear mixed effects model was used to analyse overall hand 201 pain NRS over time. The model assumed an exchangeable covariance structure to account 202 for the repeated measures over time, and included fixed effects of time (3, 6, 12-months), 203 treatment group, time-by-treatment interaction, and the pre-specified covariates (baseline 204 hand pain severity, average grip strength, concomitant analgesic use, age, gender and BMI). The model estimate of group differences at 6-months constituted the primary endpoint of the 205 206 trial. As the mixed-effects analysis model incorporated follow-up data from all available timepoints simultaneously, participants with valid outcome data at one or more follow-up visits 207 208 and complete baseline covariate data were included. Secondary analyses explored robustness to adjustments based on treatment adherence up to 6-months (binary, based on 209 self-reported non-adherence, treatment withdrawals and receipt of corticosteroids; analysis 210 using complier-average causal effect (CACE); implemented using instrumental variable 211 212 analysis (35)), 'missingness' (using multiple imputation by chained equations) and receipt of rescue medication during follow-up (increased dose or addition of any NSAIDs, opioids or 213 paracetamol or steroid injection to the hand, added as a time varying covariate (36)), all 214 detailed further in Appendix 4. The primary analysis was repeated for participants with OA 215 confirmed by imaging. To account for deviations between intended and achieved follow-up 216 timing, predicted effects at 3, 6, and 12-months were obtained from a mixed effects model, 217 218 including time of response since randomization as a continuous variable with a random 219 slope.

220

Planned sub-group analyses explored differences in treatment response for different levels
 of structural damage (mild/moderate versus severe damage based on Kallman score tertiles)
 and treatment differences in the presence/absence of ultrasound synovitis (assessed by

greyscale, Power Doppler and total synovitis) and osteophytes. Analyses were conducted by adding an interaction term between treatment allocation and the sub-groups to the primary analysis model. In the interest of planning future research, effectiveness was explored across four further sub-groups that were hypothesised to affect the treatment mechanism of HCQ, specifically average grip strength (low (<30lbs) and high strength (≥30lbs) based on median strength at baseline) and presence/absence of thumb pain.

230

Due to the large number of secondary outcomes, only outcomes of primary clinical interest were analysed using mixed-effects models, giving treatment effect estimates and p-values at each follow-up point. The remaining secondary outcomes were reported descriptively only.

235 Role of the funding source

HERO was funded by an Arthritis Research UK Clinical Studies Grant (Reference 19545).

237 Arthritis Research UK were not involved in the study design, conduct, analysis, data

interpretation, manuscript preparation or decision to submit the manuscript for publication.

239

240 **Results**

Of 316 patients screened, the HERO trial recruited 248 participants (74.5%, 124 in each trial 241 arm) with hand OA from 13 centres in England, while 68 patients were excluded (Appendix 242 243 Figure 1). Baseline characteristics (Table 1) were balanced across treatment arms. 244 Participants were on average 62.7 years old (SD=9.1), 81.9% women, predominantly of Caucasian ethnicity and had been suffering with hand pain for a median of 5 years. Nearly 245 all participants (89.9%) were taking analgesic medication for their hand OA, and median 246 247 hand pain over the past two weeks was 7 points on the 0 to 10 NRS. Five participants had raised Rheumatoid Factor (RF) and one had raised anti-cyclic citrullinated peptide (CCP). In 248 all six cases this was determined to be non-clinically significant by the site PI and not 249 indicative of inflammatory arthritis. 250

Most participants (70.6%) were prescribed a 300 mg daily dose of investigational medicinal 252 product (IMP, HCQ: 85, placebo: 90, Appendix Table 2), with all but one participant 253 remaining on the same dose throughout the trial. Balance in participant characteristics was 254 255 maintained for patients included in the intention-to-treat analysis. In total, 45 participants 256 (18.1%, HCQ: 24, placebo: 21) were non-adherent to the treatment, which is likely to be a conservative estimate, assuming unknown, unreported non-adherence. Non-adherers 257 258 tended to be slightly younger (mean of 61.2 years versus 63.0 years) with greater average 259 grip strength (36.1lbs versus 31.3lbs). Follow-up was 84.7% at 6-months and 76.6% at 12months. A total of 134 participants (54.0%) received rescue medication during the trial 260 261 (HCQ: 63, placebo: 71).

262

263 Primary Outcome

Hand pain severity improved for participants with observed data in both arms by around 1 264 point between baseline and 3 months, and this was maintained up to 12-months (Figure 1A). 265 Outcome data was not available for 20 patients at 3-months, 38 patients at 6-months and 58 266 267 patients at 12-months follow-up (Appendix Figure 1). A total of 232 participants (93.5%, HCQ: 113, placebo:119) were included in the primary intention-to-treat analysis. Differences 268 in hand pain severity between treatment groups were small at each follow-up and not 269 statistically significant (Table 2; Figure 1A). At the 6-month primary endpoint, the treatment 270 difference estimate was -0.16 points on the NRS pain scale (95% CI: -0.73 to 0.40, p=0.57), 271 i.e. participants in the HCQ arm reported worse pain by 0.16 score points, equivalent to a 272 standard effect size of 0.07. The confidence interval excludes a clinically meaningful 273 difference in improvement of 0.8 scale points, on which the trial was powered. Improvements 274 of this magnitude or greater were reported for 58 of 107 patients in the HCQ group and 59 of 275 276 103 patients in the placebo group with NRS pain score reported at 6-months.

277

278 Results were robust to secondary analyses of hand pain severity. When non-adherence was accounted for, the treatment effect became positive (0.21 scale points in favour of HCQ). 279 While the 95% confidence interval remained wide (-0.44 to 0.86), the upper limit did include 280 281 the potentially meaningful clinical difference of 0.8 scale points (Table 2). When multiple 282 imputation was used to address missing outcome and baseline grip strength data, results 283 were comparable with the primary analysis of hand pain severity with similar confidence 284 interval widths (Table 2). Treatment effects of the analysis accounting for rescue medication 285 closely resembled those of the primary analysis of hand pain severity (Table 2). A repeat 286 analysis for participants with confirmed OA on imaging (n=171 of 182 with available imaging data and analysis covariates) as well as estimates treating response time continuously 287 288 revealed no significant treatment differences (Appendix Table 3), with confidence intervals 289 excluding a clinically meaningful difference.

290

291 Safety

A total of 15 serious adverse events (SAEs) were reported by 15 patients (HCQ: 7, placebo:

8; Appendix Table 5). No deaths were reported. Of the 15 SAEs, three were assessed as

being related to HCQ: prolonged QT interval with ventricular arrhythmias, erythema

295 multiforme and acute generalised erythematous pustulosis.

296

297 Secondary Outcomes, Subgroup Analyses and Ultrasound Findings

Hand pain and most self-reported symptom outcomes improved in the short term in both arms
and then plateaued over follow-up. Mental functioning outcomes, grip strength and structural
damage remained unchanged. There were no systematic treatment differences between HCQ
and placebo for any of the secondary outcomes (Table 3, Appendix Table 4). A difference of
borderline statistical significance (SF-12 physical component score at 12 months (p=0.053))
could be spurious in light of the number of outcomes and timepoints assessed.

304

305 Radiograph data at baseline, recorded as Kallman scores, were available for 188 participants (75.8%), 94 in each arm. Data tertiles were used to group observations into mild 306 307 to moderate damage (score 0-57) and severe damage (score 58-113). There were no 308 substantial differences between severity groups in response to treatment, and the value of a 309 group by treatment interaction term added to the primary analysis model was not statistically 310 significant (p=0.25; Figure 1B). A significant interaction term with treatment allocation 311 (p=0.033) indicated that participants with greater grip strength may benefit more from HCQ 312 treatment than weaker participants (Appendix Figure 2). A treatment interaction with 313 baseline thumb pain did not reveal meaningful group differences (p=0.136, Appendix Figure 3). As the latter two analyses were exploratory, results may be considered spurious. 314 315

Baseline ultrasound images were taken for a subset of randomized participants (n=143, 57.7%; HCQ: 74, placebo: 67). The vast majority were positive for synovitis assessed by greyscale (93.7%) and over half for synovitis assessed by Power Doppler (58.7%). Osteophytes were present in at least one joint for all participants. There were no significant treatment differences between participants with positive or negative Power Doppler status

(p=0.85 for the interaction term with treatment, Figure 1C). Meaningful sub-group analyses
 were not possible for greyscale synovitis (only nine negative cases), total synovitis (Power
 Doppler did not add new cases) or osteophytes.

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325 Conclusions

The HERO trial was designed as a pragmatic trial with a view to replicating anecdotal reports of HCQ use in clinical practice, and powered to detect a moderate effect equivalent to that for NSAIDs in this population. We found that HCQ was not a more effective analgesic than placebo when added to usual care in people with moderate to severe hand OA. There were no demographic differences in the patient population that might explain the lack of efficacy. Background analgesic use did not differ between groups and baseline inflammation and structural damage did not affect response to HCQ. The study therefore presents no evidence

to suggest that HCQ should be considered within the management plan of people with handOA.

335

In terms of age, gender distribution and BMI, our population reflects that observed in recent 336 337 community-based cohorts of hand OA in the UK and Europe (37-40). We deliberately excluded participants with isolated 1st carpometacarpal joint (CMCJ) involvement or 338 predominant 1st CMCJ pain, due to the potential differences in mechanism of disease 339 340 between 1st CMCJ and distal and proximal interphalangeal joint OA. Whilst just over half of 341 participants had concomitant thumb pain, in line with previous community studies (37-40), this was not the primary site of their hand pain and no difference in treatment effect was 342 observed in those with or without CMCJ involvement. Consistent with recent imaging 343 studies, ultrasound-detected greyscale synovitis was common, with nearly all participants 344 345 having moderate grade synovitis in at least one joint. Power Doppler synovitis although less common, present in just over half of participants, was not associated with treatment 346 differences. Based on the additional sub-group analyses, weaker grip strength may 347 predispose people to tenosynovitis or enthesitis, alternative causes for hand pain in this 348 349 population. This suggests a need to consider grip strength in this population when planning 350 further studies.

351

A growing body of imaging and experimental evidence suggests a role for synovitis in the 352 pathogenesis of OA and an association with pain. Ultrasound-detected synovitis is 353 independently associated with radiographic progression of hand OA, painful hand joints are 354 associated with the presence of ultrasound- and MRI-detected synovitis, and response to 355 356 intramuscular steroids (thought to work by reducing synovitis) in hand OA is associated with higher levels of baseline ultrasound-detected synovitis (19,41-44). However, in the HERO 357 study baseline synovitis was not linked to treatment effect. Our inclusion criteria may have 358 359 resulted in participants where the level and/or type of inflammation was not severe: a 360 previous study has suggested that early OA may be more inflammatory than established OA,

and that molecular pathways driving inflammation may change as the disease progresses
(45). By selecting participants with moderate to severe hand OA, established radiographic
changes and inadequate response to existing therapies, we may have missed an early
window of opportunity for HCQ to have therapeutic benefit.

365

366 Hydroxychloroquine has various known immunomodulatory effects, and although 367 established as a treatment option in the management of inflammatory arthritides, its specific 368 mechanism of action remains unclear. In RA, therapeutic activity has been linked to 369 modulation of antigen-processing activity, including inhibition of T-cell activation and cytokine 370 release (46,47); increasing evidence of involvement of these pathways in inflammation and 371 cartilage degeneration in OA (48-50) supported HCQ as a potential OA therapy. More recent data implicates intracellular toll-like receptors (TLR), in particular TLR-9, as key mediators of 372 373 HCQ's anti-inflammatory properties, in line with growing evidence of the role of the innate immune system in rheumatic disease. Although limited evidence suggests that the innate 374 immune system may be important in OA pathogenesis (51), for example increased TLR 375 expression in OA tissue (52-55), this work is still in its infancy. Further understanding of 376 377 these mechanisms in OA may enable stratification according to a defined inflammatory 378 phenotype.

379

Other potential limitations to the study include restriction of HCQ dosing to the British 380 National Formulary recommended maximum dose of 6.5 mg/kg/day (56), with the majority of 381 patients taking 300 mg daily. In clinical RA practice, patients may commence HCQ at a 382 higher dose (400 mg), with reduction to a lower maintenance dose after 3-6 months. 383 384 However, only 5.6% of the HCQ group were on the lowest dose of 200mg and no dose-385 response relationship with treatment effect was observed. The co-occurrence of MRIdetected bone marrow lesions (BMLs) with hand synovitis has been found to worsen pain 386 387 and, as demonstrated in knee OA, may contribute to pain (57,58). Since BMLs cannot be 388 detected by ultrasound or x-ray, we were unable to examine BMLs in this study. The failure

389 of HCQ as an analgesic in this study may reflect the mild anti-inflammatory activity of HCQ, suboptimal dosing, or that the level and/or type of inflammation in our population did not 390 match the mechanism of HCQ. However it is also worth considering, in light of the current 391 result and the previous failure of biologic DMARDs, that simply treating 'macroscopic' or 392 393 imaging-detected synovitis with DMARDs is not a useful analgesic strategy. Further 394 exploration of the molecular mechanisms of inflammation in OA may provide targets and 395 better patient phenotyping may enable exclusion of other causes of hand pain such as 396 tenosynovitis.

397

In summary, HCQ was not more effective than placebo in reducing symptoms or
radiographic progression in people selected for moderate to severe hand pain and
radiographic OA. Our findings in this full-scale pragmatic trial do not support the current
practice for the off-label use of Hydroxychloroquine in those with hand osteoarthritis.

402

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410

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419

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586

587 Figure Legends

- 588 Figure 1: Unadjusted Hand Pain NRS (past two weeks) with 95% CIs; A) HERO study
- 589 participants with observed data (primary outcome). B) Structural damage sub-groups (based

590	on Kallman total score); C) Synovitis sub-groups (ultrasound sub-study). HCQ =
591	hydroxychloroquine.
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647	15 Cannock Chase Hospital, Brunswick Road, Cannock, WS11 5XY, UK.
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Table 1: Baseline Characteristics

HCQ Placebo mmary analysis (n=232) Age (n=124) (n=124) HCQ Placebo N 124 124 113 119 Mean (SD) 62.8 (9.1) 62.5 (9.2) 63.1 (9.3) 62 (40.83) Gender 1 64 (41,88) 62 (40.83) 64 (41,88) 62 (40.83) Gender 2 7 (27%) 106 (85%) 87 (77%) 102 (86%) BMI 7 124 124 113 119 Mean (SD) 28.4 (5.4) 29.3 (6.2) 28.5 (5.4) 29.4 (6.3) Median (min, max) 28 (15, 45) 28 (15, 45) 28 (19, 45) 28 (19, 45) Ethnicity 119 (96%) 120 (97%) 109 (96%) 116 (97%) South Asian 1 (1%) 1 (1%) 1 (1%) 1 (1%) 1 (1%) Caucasian 119 (96%) 120 (97%) 109 (96%) 116 (97%) South Asian 1 (1%) 1 (1%) 1 (1%) 1 (1%) Median (min, max) 2 (2%) 1 (1%)		All randomised patients		All patients included in the		
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Other 1 (1%) 2 (2%) 0 (0%) 1 (1%) Hand pain duration in years 124 124 113 119 Mean (SD) 7.4 (6.4) 7.9 (6.7) 7.7 (6.5) 7.8 (6.8) Median (min, max) 5 (0.4, 30) 5.5 (1, 30) 6 (0.4, 30) 5.5 (1, 30) Hand Pain NRS (past 48 hours) [0 none - 10 worst] 113 117 N 124 121 113 117 Mean (SD) 6.9 (1.7) 6.8 (1.8) 6.9 (1.62) 6.8 (1.77) Median (min, max) 7 (2, 10) 7 (2, 10) 7 (3, 10) 7 (2, 10) Grip Strength in lbs (average both hands) 29.9 (19.3) 34.6 (19.6) 29.4 (18.9) Median (min, max) 31.3 (0, 114.2) 27.5 (1.0, 95.0) 95.0) 95.0) AUSCAN Pain [0-20] 12.4 121 113 117 Mean (SD) 12.3 (2.61) 12.7 (3.00) 12.4 (2.6) 12.7 (3.0) Median (min, max) 12.5 (4, 18) 13 (4, 20) 13 (4, 18) 13 (4, 20) N 12.3 (2.61) <	Afro-Caribbean	1 (1%)	0 (0%)	1 (1%)	0 (0%)	
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Median (min, max) 7 (0, 33) 8 (0, 38) 7 (0, 33) 7 (0, 38) Total number of painful joints [0-30] 124 124 113 119 Mean (SD) 8.3 (5.9) 8.8 (7.1) 8.5 (5.9) 8.6 (7.0) Median (min, max) 7 (0, 30) 7 (0, 30) 6 (0, 30)	Mean (SD)	95(95)	10.8 (9.5)	98(96)	10.5 (9.5)	
Total number of painful joints 1 (0, 00) 1 (0, 00) 1 (0, 00) N 124 124 113 119 Mean (SD) 8.3 (5.9) 8.8 (7.1) 8.5 (5.9) 8.6 (7.0) Median (min, max) 7 (0, 30) 7 (0, 30) 7 (0, 30) 6 (0, 30)	Median (min_max)	7 (0, 33)	8 (0, 38)	7 (0, 33)	7 (0, 38)	
[0-30] 124 124 113 119 Mean (SD) 8.3 (5.9) 8.8 (7.1) 8.5 (5.9) 8.6 (7.0) Median (min, max) 7 (0, 30) 7 (0, 30) 7 (0, 30) 6 (0, 30)	Total number of painful joints		- (0, 00)	. (0, 00)		
N 124 124 113 119 Mean (SD) 8.3 (5.9) 8.8 (7.1) 8.5 (5.9) 8.6 (7.0) Median (min, max) 7 (0, 30) 7 (0, 30) 7 (0, 30) 6 (0, 30)	[0-30]					
Mean (SD) 8.3 (5.9) 8.8 (7.1) 8.5 (5.9) 8.6 (7.0) Median (min, max) 7 (0, 30) 7 (0, 30) 7 (0, 30) 6 (0, 30)	N	124	124	113	119	
Median (min, max) $7(0, 30)$ $7(0, 30)$ $7(0, 30)$ $6(0, 30)$	Mean (SD)	8.3 (5.9)	8.8 (7.1)	8.5 (5.9)	8.6 (7.0)	
	Median (min, max)	7 (0, 30)	7 (0, 30)	7 (0, 30)	6 (0, 30)	

	All randomised patients		All patients included in the	
	(n=248)		primary analysis (n=232)	
	HCQ	Placebo	HCQ	Placebo
	(n=124)	(n=124)	(n=113)	(n=119)
umber of swollen joints [0-				
30]				
N	124	124	113	119
Mean (SD)	3.8 (4.2)	3.4 (4.4)	4.0 (4.3)	3.4 (4.4)
Median (min, max)	3 (0, 20)	1 (0, 22)	3 (0, 20)	1 (0, 22)
umber of tender joints [0-30]				
N	124	124	113	119
Mean (SD)	10.4 (6.3)	10.9 (7.3)	10.4 (6.3)	10.8 (7.3)
Median (min, max)	10 (0, 27)	9 (0, 30)	10 (0, 27)	9 (0, 30)
Pain in other joints present	114 (92%)	107 (86%)	103 (91%)	102 (86%)
Number of other painful				
joints [0-14]				
N	124	123	113	119
Mean (SD)	5.8 (2.8)	5.9 (3.1)	5.9 (2.7)	5.8 (3.0)
Median (min, max)	6 (0, 12)	5 (0, 14)	6 (0, 12)	5 (1, 14)
Kallman total radiograph				
score				
N	94	94	89	93
Mean (SD)	42.7 (25.9)	47.2 (27.4)	43.9 (25.8)	47.3 (27.5)
Median (min, max)	40 (0, 100)	39 (2, 113)	41 (0, 100)	40 (2, 113)
Medication for hand OA				
Oral NSAIDs	50 (40%)	53 (43%)	49 (43%)	50 (42%)
Topical NSAIDs	22 (18%)	25 (20%)	22 (19%)	23 (19%)
Paracetamol	77 (62%)	75 (60%)	69 (61%)	70 (60%)
Opioids	14 (11%)	16 (13%)	12 (11%)	14 (12%)
Co-codamol	23 (19%)	26 (21%)	22 (19%)	26 (22%)
Other	15 (12%)	20 (16%)	14 (12%)	19 (16%)
Any concomitant analgesic	111 (90%)	112 (90%)	101 (89%)	107 (90%)
use				
Currently using glucosamine and/or chondroitin	20 (16%)	17 (14%)	19 (17%)	15 (13%)

AUSCAN = Australian/Canadian Hand Osteoarthritis Index; BMI = body mass index; HCQ =

hydroxychloroquine; NRS = numerical rating scale; NSAIDs = non-selective anti-inflammatory drugs;

OAQoL = Osteoarthritis Quality of Life

Analysis & Follow-up	N	HCQ Mean (95% CI)	N	Placebo Mean (95% Cl)	Difference Mean (95% CI)	p-value			
Primary Analy	ysis †								
3 months	113	5.54 (5.01, 6.07)	119	5.78 (5.26, 6.29)	0.24 (-0.31, 0.78)	.40			
6 months *	113	5.66 (5.13, 6.19)	119	5.49 (4.96, 6.02)	-0.16 (-0.73, 0.40)	.57			
12 months	113	5.39 (4.83, 5.92)	119	5.51 (4.98, 6.04)	0.13 (-0.45, 0.72)	.66			
Adherence ad	Adherence adjusted analysis (CACE) ‡								
6 months	107	5.53 (5.12, 5.94)	103	5.74 (5.29, 6.19)	0.21 (-0.44, 0.86)	.52			
Analysis inclu	uding a	II randomized particip	pants u	sing multiple imputat	ion §				
3 months	124	5.53 (4.98, 6.08)	124	5.76 (5.22, 6.30)	0.23 (-0.31, 0.78)	.40			
6 months	124	5.65 (5.11, 6.18)	124	5.45 (4.89, 6.00)	-0.20 (-0.80, 0.41)	.52			
12 months	124	5.38 (4.79, 5.97)	124	5.55 (5.02, 6.08)	0.17 (-0.43, 0.77)	.58			
Analysis adjusted for receipt of rescue medication									
3 months	113	5.63 (5.09, 6.17)	119	5.87 (5.34, 6.39)	0.23 (-0.31, 0.78)	.40			
6 months	113	5.70 (5.16, 6.23)	119	5.52 (4.99, 6.05)	-0.18 (-0.74, 0.38)	.53			
12 months	113	5.36 (4.82, 5.91)	119	5.48 (4.95, 6.01)	0.12 (-0.47, 0.70)	.69			

Table 2: Estimated Treatment Differences in Mean Hand Pain NRS (last 2 weeks)

* Primary Endpoint

† Linear mixed effects model with fixed effects of treatment, time, treatment by time interaction, baseline hand pain, age, gender, BMI, baseline grip strength and baseline concomitant analgesic use

‡ Instrumental variable regression(35; Appendix 5) of the outcome at 6 months, accounting for adherence with the active treatment, baseline hand pain, age, gender, BMI, baseline grip strength and baseline concomitant analgesic use

§ Linear mixed effects model with fixed effects of treatment, time, treatment by time interaction, baseline hand pain, age, gender, BMI, baseline grip strength and baseline concomitant analgesic use (any missing data was imputed from analysis covariates using multiple imputation by chained equations) (Appendix 5)

|| Linear mixed effects model with fixed effects of treatment, time, treatment by time interaction, baseline hand pain, age, gender, BMI, baseline grip strength and baseline concomitant analgesic use and receipt of rescue medication (time varying) (REF: White et al, 2001; Appendix 5)

HCQ = hydroxychloroquine; NRS = numerical rating scale measured using an 11-point (0-10) scale;

Outcome & Follow-up	N	HCQ Mean (95% CI)	N	Placebo Mean (95% Cl)	Difference Mean (95% CI)	p-value	
Pain severity	in the r	nost painful joint (NR	S over	last 2 weeks, range 0-1	0, higher score = worse	pain) *	
3 months	112	5.85 (5.31, 6.40)	119	5.49 (4.96, 6.02)	0.19 (-0.37, 0.75)	.51	
6 months	112	6.20 (5.66, 6.75)	119	5.85 (5.31, 6.40)	-0.30 (-0.88, 0.28)	.31	
12 months	112	5.83 (5.27, 6.40)	119	6.20 (5.66, 6.75)	-0.09 (-0.70, 0.51)	.76	
AUSCAN Pair	ı (Rang	e: 0-20, higher score =	worse	functioning) †		•	
3 months	113	11.29 (10.48, 12.11)	117	11.22 (10.42, 12.02)	-0.07 (-0.91, 0.77)	.87	
6 months	113	11.14 (10.32, 11.96)	117	10.99 (10.17, 11.81)	-0.15 (-1.02, 0.71)	.73	
12 months	113	10.92 (10.08, 11.76)	117	10.38 (9.55, 11.20)	-0.55 (1.44, 0.35)	.23	
AUSCAN Fun	ction (F	Range: 0-36, higher sco	ore = wo	orse functioning) ‡			
3 months	112	19.61 (18.19, 21.03)	118	20.04 (18.64, 21.43)	0.43 (-1.05, 1.90)	.57	
6 months	112	19.51 (18.07, 20.94)	118	19.19 (17.76, 20.61)	-0.32 (-1.84, 1.20)	.68	
12 months	112	19.72 (18.24, 21.20)	118	18.74 (17.30, 20.18)	-0.98 (-2.55, 0.59)	.22	
Grip Strength	Left Ha	and (in lbs) §					
6 months	105	36.95 (33.26, 40.64)	104	37.98 (34.31, 41.65)	1.03 (-2.75, 4.82)	.59	
12 months	105	37.08 (33.31, 40.85)	104	38.85 (35.12, 42.58)	1.77 (-2.14, 5.68)	.38	
Grip Strength	Right	Hand (in lbs) §					
6 months	105	37.34 (33.71, 40.97)	103	37.25 (33.63, 40.88)	-0.09 (-3.87, 3.69)	.96	
12 months	105	36.79 (33.08, 40.50)	103	38.89 (35.24, 42.54)	2.10 (-1.80, 5.99)	.29	
Kallman Tota	l Radio	graph Score (Range:	0-220, h	igher score = greater s	structural damage)		
12 months	79	48.14 (47.32, 48.96)	78	48.30 (47.50, 49.10)	0.16 (-0.69, 1.00)	.72	
Osteoarthritis	s Qualit	y of Life (OAQol, rang	e: 0-38,	higher score = greater	impact of OA symptoms	s) ¶	
6 months	106	8.60 (7.25, 9.95)	102	8.83 (7.50, 10.17)	0.24 (-1.13, 1.60)	.74	
12 months	106	8.96 (7.58, 10.35)	102	9.58 (8.23, 10.94)	0.62 (-0.80, 2.05)	.39	
SF-12 Physic	al Com	ponent Score (Range:	: 0-100,	higher score = better f	unctioning) **		
6 months	107	39.63 (37.50, 41.77)	104	39.70 (37.57, 41.82)	0.07 (-2.14, 2.28)	.95	
12 months	107	38.32 (36.11, 40.53)	104	40.58 (38.44, 42.72)	2.26 (-0.03, 4.55)	.053	
SF-12 Mental Component Score (Range: 0-100, higher score = better functioning) ++							
6 months	107	51.52 (49.34, 53.69)	104	52.24 (50.09, 54.38)	0.72 (-1.57, 3.01)	.54	
12 months	107	53.15 (50.89, 55.40)	104	52.00 (49.83, 54.17)	-1.15 (-3.53, 1.24)	.35	

Table 3: Key Secondary Outcomes - Mean Estimates from Analysis Models

* Linear mixed effects model with fixed effects of treatment, time, treatment by time interaction, baseline pain severity, age, gender, BMI, baseline grip strength and baseline concomitant analgesic use

† Linear mixed effects model with fixed effects of treatment, time, treatment by time interaction, baseline AUSCAN pain, age, gender, BMI, baseline grip strength and baseline concomitant analgesic use

‡ Linear mixed effects model with fixed effects of treatment, time, treatment by time interaction, baseline AUSCAN function, age, gender, BMI, baseline grip strength and baseline concomitant analgesic use

§ Linear mixed effects model with fixed effects of treatment, time, treatment by time interaction, baseline grip strength, age, gender, BMI and baseline concomitant analgesic use

|| Linear regression model with fixed effects of treatment, baseline Kallman score, age, gender, BMI, baseline grip strength and baseline concomitant analgesic use

I Linear mixed effects model with fixed effects of treatment, time, treatment by time interaction, baseline OAQoI, age, gender, BMI, baseline grip strength and baseline concomitant analgesic use

** Linear mixed effects model with fixed effects of treatment, time, treatment by time interaction, baseline SF-12 PCS, age, gender, BMI, baseline grip strength and baseline concomitant analgesic use

†† Linear mixed effects model with fixed effects of treatment, time and treatment by time interaction, adjusted for baseline SF-12 MCS, age, gender, BMI, baseline grip strength and baseline concomitant analgesic use

AUSCAN = Australian/Canadian Hand Osteoarthritis Index; NRS = numerical rating scale; OAQoL = Osteoarthritis Quality of Life; SF-12 = Short Form - 12



Figure 1: Unadjusted Hand Pain NRS (past two weeks) with 95% CIs

