Chloroquine and Hydroxychloroquine Retinal Toxicity Consideration in the Treatment of COVID-19

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Abstract: The proposed doses of chloroquine (CQ) and hydroxychloroquine (HCQ) for treatment of COVID-19 (1000 mg/day for 10 days, CQ; 800 mg first day then 400 mg/day for 5 days, HCQ) in many guidelines worldwide, are considerably higher than the maximum recommended daily safe doses of both agents (≤2.3 mg/kg/day, CQ; ≤5.0 mg/kg/day, HCQ) for development of retinal toxicity. Irreversible retinal damage can occur if exposure to the safe doses is ≥5 years. It is not known whether exposure to high doses over a short period of time can also cause the damage. We recommend that before prescribing CQ or HCQ, history of ocular disease should be obtained to avoid the prescription if appropriate. If either agent is to be used, routine baseline ocular examination is not absolutely necessary. Patients who do not have ocular disease should also be informed about the potential risk of retinal toxicity. Both agents, however, have not yet been proven to be beneficial to COVID-19.

Key Words: chloroquine, coronavirus, hydroxychloroquine, maculopathy, retinopathy

With the occurrence of pandemic of the coronavirus disease 2019 (COVID-19) announced by the World Health Organization in early March 2020 and the number of cases still on the rise in all continents in late March, many therapeutic options have been proposed for this novel and potentially fatal disease. Apart from antiviral agents, chloroquine (CQ) and hydroxychloroquine (HCQ) have been examined for their roles in treatment of COVID-19. This may be because both CQ and HCQ have been postulated to reduce viral replication in other coronavirus infections. According to a recent systematic review, there are almost 20 ongoing randomized controlled clinical trials on both medications for treatment of COVID-19 and all of them are in China. The details including dosing regimens of CQ and HCQ in these trials have been summarized in the review. Although there has not yet been a completed clinical trial and the world is waiting eagerly for the results of these trials and other trials on other treatment options, many authorities have chosen to adopt CQ and HCQ in the guidelines for treatment of COVID-19 based on in vitro studies, nonrandomized trial, and anecdotal evidence. As the therapeutic doses of CQ and HCQ recommended in the trials and guidelines are relatively high compared with the maximum daily safe dose that is related to CQ and HCQ retinal toxicity, this issue of retinal toxicity should be taken into consideration when employing these 2 medications for treatment of COVID-19 worldwide.

According to the recommendation by the American Academy of Ophthalmology, the most significant major risk factors for CQ and HCQ retinal toxicity are high dose and long duration of use. Other risk factors include concomitant renal disease and use of tamoxifen. The maximum daily dose from this recommendation is ≤5.0 mg/kg real body weight for HCQ, and ≤2.3 mg/kg real body weight for CQ. As shown in Table 1, the doses of CQ or HCQ for treatment of COVID-19 in various treatment guidelines worldwide are well beyond these recommended dosing regimens.

The Royal College of Ophthalmologists (RCO) in the UK also addressed the importance of safe dose and duration of prescription of CQ and HCQ for the development of retinal toxicity. Although no absolute safe dose was identified, the RCO recommends the daily dose of HCQ to be <5 mg/kg/day for <5 years as relatively safe for retinal toxicity. However, no safe dose of CQ was recommended and the RCO identified those who receive CQ for >1 year as having risk of retinal toxicity.

Despite the fact that the daily doses of both CQ and HCQ for the treatment of COVID-19 exceed the daily safe doses of both agents, the treatment may still be considered relatively safe for retinal toxicity. The toxicity, which causes irreversible retinal damage and visual loss despite ceasing prescription, requires exposure to the safe dose for a long period of time, generally in excess of 5 years. In general, the recommendation for screening for retinal toxicity from CQ and HCQ is within the first year of use as baseline and then annual screening after a year of use for CQ and 5 years of use for HCQ. Both American Academy of Ophthalmology and RCO recommended the screening should be conducted sooner if the major risk factors are present.

In the case of treatment of COVID-19 using CQ and HCQ, the
Table 1. Comparison Between the Maximum Daily Safe Doses of Chloroquine and Hydroxychloroquine for Development of Retinal Toxicity and the Recommended Doses in COVID-19 Treatment Guidelines From Different Countries

| Country/Guideline | Maximum Recommended Dosing Regimen | Daily Safe Dose for Retinal Toxicity |
|-------------------|-----------------------------------|-------------------------------------|
| Dutch Guideline7   | 1000 mg/day for at least 5 days    | 2.3 mg/kg/day estimated global dose  |
| Belgian Guideline9 | 1000 mg/day for up to 5 days       | 600 mg Day 1, then 800 mg Day 1, then 400 mg/day up to 5 days |
| Italian Guideline11| 800 mg Day 1, then 800 mg Day 1, then 400 mg/day up to 5 days |
| Chinese Guideline8 | 800 mg Day 1, then 800 mg Day 1, then 400 mg/day up to 5 days |
| Thai Guideline6    | N/A 800 mg Day 1, then 400 mg/day up to 5 days |

The estimated global doses are estimated according to the global average weight of adults (62 kg).17

In summary, the bottom line at the present time is that neither CQ nor HCQ has been proven to be effective in the treatment of COVID-19, although there is certainly a vast interest in its possible benefit. Further controlled clinical trial data will be necessary to help better address this issue. Despite the current situation of COVID-19 pandemic, many adverse effects of either CQ or HCQ should still be weighed against its potential benefit. For retinal toxicity, the risk of having irreversible retinal damage.
and visual loss may outweigh the unproven benefit of both agents in some patients. Detecting the risk is easy. It can be done by simply taking a history of previous or co-existing ocular disease from the patients, then other options of treatment should be considered if appropriate.

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