Stopping prer eferral rectal artesunate — a grave error

James A Watson,1,2 Marian Warsame,3 Thomas J Peto,1 Marie Onyamboko,4,5 Caterina Fanello,1,2 Arjen M Dondorp,1,2 Nicholas White1,2

INTRODUCTION

It is estimated that over half a million people, mainly African children, died last year of severe Plasmodium falciparum malaria.1 A key pathogenic process in severe falciparum malaria is the unchecked intravascular multiplication of malaria parasites to reach burdens which cause potentially lethal microvascular obstruction in vital organs. Severe malaria is rapidly fatal if untreated.2 Artemisinins are the cornerstone of management.3,4 There is overwhelming evidence that if adequate concentrations of these drugs are reached in blood, then therapeutic responses are accelerated compared with other antimalarials. Parenteral artesunate saves 80%–90% of treated patients with strictly defined severe malaria, and it reduces their mortality by about one-third when compared with parenteral quinine, the previous first-line treatment.2,5

However, in many rural settings, parenteral drug administration is not possible.2,6 Following pioneering studies, first in China, and then in Vietnam, the rectal route of administration for artemisinin was shown to provide a simple prer eferral approach in rural settings to the management of patients who could not retain oral antimalarial medicines, or who had suspected severe malaria.2,6–11 Between August 2000 and July 2006, a very large double-blind placebo-controlled randomised trial of a pre-referral rectal formulation of artesunate (RAS) in patients with suspected severe malaria was conducted in Tanzania, Ghana and Bangladesh.6 The overall mortality was 2.5% (154/6072) in the placebo arm, 3.0% (177/5996; p=0.1) in the placebo arm, although many of the studied patients did not have malaria. These case fatality rates were much lower than expected when this trial was designed, probably because of the support and transport provided for the community health workers (CHWs). In a post hoc subgroup analysis of the patients who did have malaria, but took more than 6 hours after the administration of rectal artesunate to reach the referral centre for parenteral therapy, 1.9% (29/1566) of patients in the RAS arm died or had permanent disabilities compared with 3.8% (57/1519) in the placebo arm (p=0.0013).

Meanwhile, large randomised controlled trials of hospitalised patients in Asia and Africa showed that parenteral artesunate reduced the mortality of severe falciparum malaria by 35% and 22.5%, respectively.12 Rectal artesunate, therefore, solved the problem in remote rural areas of potentially lethal delays in receiving appropriate treatment. A life-saving strategy for severe malaria is needed now more than ever.2,13

Summary box

⇒ The WHO recently recommended a moratorium on the implementation of pre-referral rectal artesunate for suspected childhood severe malaria.
⇒ This was because of the lower referral completion rates and higher case fatality ratios in rectal artesunate recipients reported in a non-randomised, sequential observational study (CARAMAL) conducted in Nigeria, Uganda and the Democratic Republic of Congo.
⇒ The observational study design, lack of a prespecified statistical analysis plan, strong temporal confounding, likely selection bias and the biological implausibility, all strongly challenge the interpretation of a causal relationship between rectal artesunate roll-out and increased mortality.
⇒ Early mortality associated with delayed referral following rectal artesunate administration is more likely to have resulted from inadequate treatment of sepsis rather than malaria.
⇒ Further delays in rectal artesunate roll-out resulting from inappropriate analysis and interpretation of the CARAMAL study will likely result in preventable childhood mortality.
⇒ Stopping prer eferral rectal artesunate is a grave error. The WHO should lift its moratorium on rectal artesunate deployment without delay.
antimalarial could now be given in the village to children with suspected malaria who could not swallow oral medicines reliably. Unfortunately, the subsequent deployment of rectal artesunate has gone very slowly, and now it has been halted.

**THE CARAMAL STUDY**

This policy reverse is because of a large, ‘real world’, sequential observational study (CARAMAL), which evaluated the implementation of RAS in the Democratic Republic of Congo (DRC), Uganda and Nigeria. The study described shortcomings in patient referral to health facilities after they had received RAS, and incomplete follow-up treatment with oral artemisinin combination treatments. CARAMAL provides useful guidance for improving the deployment of RAS. However, the study also associated the deployment and use of rectal artesunate with an apparent increase in mortality. This also associated the deployment and use of rectal artesunate with an apparent increase in mortality. This policy reverse is because of a large, ‘real world’, sequential observational study (CARAMAL), which evaluated the implementation of RAS in the Democratic Republic of Congo (DRC), Uganda and Nigeria. The study described shortcomings in patient referral to health facilities after they had received RAS, and incomplete follow-up treatment with oral artemisinin combination treatments. CARAMAL provides useful guidance for improving the deployment of RAS. However, the study also associated the deployment and use of rectal artesunate with an apparent increase in mortality.

**THE MAIN CONCERNS**

First, the sequential observational nature of the CARAMAL study makes any causal estimates of the effects of RAS roll-out unreliable. There is a potentially large bias (ie, confounding) by indication: CHWs naturally allocate potentially life-saving medicine to those who they think need it the most. These are the children who are most severely ill, and most likely to die. Substantial temporal confounding is also likely in the CARAMAL study because the periods compared and referral processes were different. The pre roll-out phase in Nigeria, where mortality was substantially higher than in DRC or Uganda, covered only half the rainy season (second half of 2018) and recruited predominantly via CHWs, whereas the major increase in mortality after roll-out occurred during the COVID-19 pandemic in patients recruited at primary health centres (PHCs). Parents may have been more reluctant to bring their sick children to healthcare facilities when there was a perceived risk of catching COVID-19. In the DRC, we have observed substantial changes in health seeking behaviour during the COVID-19 pandemic.

The CARAMAL study results were extremely heterogeneous; mortalities exceeded 10% in Nigeria but were less than 1% in Uganda. Nigeria was the only country with a significant increase in mortality after RAS roll-out (a four-fold increase: 16.1% vs 4.2%) and where, after roll-out, children receiving RAS were much more likely to die than those who did not receive RAS (2.5-fold difference: 19.7% vs 7.7%). Yet in Nigeria RAS roll-out was incomplete; only half of the Nigerian patients actually received RAS. In the 48 hours following enrolment, mortality in Nigerian patients who did receive RAS was over twice that in those who did not (10% vs 4%). This very large difference in acute mortality (within a single malaria parasite asexual life-cycle) cannot be explained by lower rates of referral and delayed administration of parenteral treatment (RAS reduces parasite numbers by 10 000-fold over one 48 hours life-cycle). Bias in RAS allocation seems a much more likely explanation, that is, healthcare providers were more likely to administer RAS to the sicker children.

Second, the authors did not write a statistical analysis plan before analysing the data. This is unusual for a large preregistered multi-country study today, and it is particularly important given that the results have changed WHO policy. Prespecifying the statistical analysis is critical to avoid data dependent analyses. There are three possible exposure contrasts (pre roll-out vs post roll-out, pre roll-out vs post roll-out and receiving RAS, and receiving vs not receiving RAS post roll-out), three different countries, and two enrolment types (CHW vs PHC), so there are at least 18 subdivisions of the data. Although the primary outcome was clearly prespecified, the reported associations were not. The lack of prespecification is illustrated by comparing the analysis of referral completion with the analysis of mortality: the set of covariates chosen for adjustment are completely different. No rationale for either set of choices was provided. This lack of prespecification means that the main reported results of the CARAMAL study, as currently presented, should be regarded as post hoc subgroup analyses.

Third, some of the key published results are misleading. They do not reflect the findings of the study. For example, in the published report on referral completion the authors state: ‘In DRC and Uganda, RAS users were less likely to complete referral than RAS non-users in the pre roll-out phase (adjusted OR (aOR)=0.48, 95% CI 0.30 to 0.77 and aOR=0.72, 95% CI 0.58 to 0.88, respectively)’. This suggests that RAS was associated with referral failure, but it misrepresents the results of actual analysis. In fact, all referral completion was lower after RAS roll-out, and it was not different in RAS non-users (indeed in DRC, patients receiving RAS were slightly more likely to complete referral than those who did not receive RAS).

Fourth, there is the pharmacological implausibility of attributing an increased proportion of rapid death to receipt of 10 mg/kg rectal artesunate. Some patients with strictly defined severe malaria do die rapidly after receiving artesunate, but over 90% survive whereas, without any treatment, nearly all die. The majority of the life-saving effect of artesunate results from the first dose. For the post-treatment burden of parasites to return to lethal levels, more than two asexual cycles (>4 days) are required. Blood stage parasites cannot reproduce more rapidly. Yet CARAMAL has been interpreted as showing that the failure to receive a second dose resulted in a substantially increased malaria-attributable mortality within 48 hours, compared with no prerelerral
treatment. A large post-RAS increase in acute malaria attributable mortality (ie, the differential) simply could not have occurred so rapidly.

Finally, we should consider the accuracy of the diagnosis of severe malaria by CHWs and PHCs. Sepsis is the major cause of preventable death in children living in malaria endemic areas. Even within prospective studies in hospital-based research centres, the diagnosis of severe malaria is incorrect in about one third of cases. The misdiagnosed children commonly have sepsis, and they have a higher case specific mortality. Twenty years ago, when the ineffective chloroquine was still the main antimalarial drug in Africa, malaria was a much more important cause of childhood death. Today, in a context of widespread insecticide treated bed-net use, access to oral artemisinin combination treatments, and increasing deployment of seasonal malaria chemoprevention, sepsis is a much more likely cause of death than malaria. Sepsis would certainly explain why delays in referral and receiving parenteral antibiotics (whether or not RAS was administered) were associated with increased mortality.

CONCLUSIONS
Prereferral RASs save lives in children with severe malaria when no parenteral treatment is available. Although recommended by WHO for years, RAS deployment has been very limited, and now it has been stopped because of the disappointing results of a non-randomised, sequential observational study (CARAMAL) conducted in Nigeria, Uganda and the DRC. The CARAMAL study was a substantial effort which showed the difficulties of ensuring timely referral of severely ill children to hospital in the three African countries. This emphasises the importance of deploying RAS in the context of sustained support for CHWs and rural health centres. But the results cannot be used to evaluate the effectiveness of rectal artesunate as a prereferral treatment of severe malaria. The observational study design cannot estimate reliably the causal effect of RAS roll-out on mortality. There was both strong temporal confounding and likely selection bias. There was no prespecified statistical analysis plan, so the many subgroup analyses reported must be considered post hoc. Furthermore, it is biologically implausible that rapid death from severe malaria would be more frequent after RAS than no treatment. The high early mortality after RAS roll-out is more likely to have resulted from sepsis than malaria.

Further delays in rectal artesunate roll-out resulting from the inappropriate analysis and interpretation of the CARAMAL study will result in preventable childhood mortality. The WHO Global Malaria Programme moratorium on RAS is unjustified, and it should be lifted without delay.

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