GRADE TABLE 2a: What is the **duration of protection/risk** in **seropositive** individuals 9-16 years of age vaccinated with CYD-TDV?

Population: 9-16 year-old individuals living in dengue endemic areas
Intervention: 3 doses of CYD-TDV administered 6 months apart
Comparison: Placebo
Outcome: Virologically-confirmed dengue occurring > 12 months of completion of 3 doses

What is the duration of protection/risk in seropositive individuals 9-16 years of age vaccinated with CYD-TDV? Rating Adjustment to rating No. of studies/starting rating 2 RCT¹ 4 Limitation in study Very serious² -1 design 0 Inconsistency None serious Factors decreasing Indirectness 0 None serious confidence 0 Imprecision None serious **Publication bias** None serious 0 **Quality Assessment** Large effect Not applicable 0 Factors Dose-response Not applicable 0 increasing confidence Antagonistic bias Not applicable 0 and confounding Final numerical rating of quality of evidence 3 Evidence supports a moderate level of confidence that the true Statement on quality of evidence effect lies close to that of the estimate of effect on Summary of Findings health outcome. CYD-TDV is associated with statistically significant protection against severe Conclusion and hospitalized dengue in seropositive participants 9-16 years of age

¹CYD-TDV has been evaluated in two parallel Phase 3 clinical trials, known as CYD14 and CYD15. CYD14 was conducted in 5 countries in Asia (Indonesia, Malaysia, Philippines, Thailand, and Vietnam), with 5,234 participants aged 9-14 years at first vaccination (10,275 participants in the full trial population aged 2-14 years). CYD15 was conducted in 5 countries in Latin America (Brazil, Colombia, Honduras, Mexico, and Puerto Rico (US)), with 20,869 participants aged 9-16 years at first vaccination. In each of these trials, participants were randomized to vaccine and placebo in a 2:1 ratio. Because the physical appearance of the vaccine and placebo was different, unmasked trial staff were responsible only for preparation and administration of injections and were not involved in the

follow-up of trial participants. For the ascertainment of trial endpoints the trials were observer-masked. Active surveillance of participants lasted only until study month 25, after which surveillance was hospital-based. Thus, it is not possible to evaluate the duration of protection against virologically-confirmed dengue of any severity. However, since severe outcomes are also of interest and importance from a public health perspective, the confidence is not downgraded.

Based on participants with serostatus derived from the new analyses, in seropositive participants aged 9-16, Hazard Ratios (HRs) up to study month 60 for hospitalized VCD and severe VCD were 0.21 (95%CI: 0.14;0.31, p<0.001) and 0.16 (95%CI: 0.07;0.37, p<0.001), respectively, and point estimates were <1 with all methods in pooled analyses and in individual trials.

²The study design of CYD14 and CYD15 included 25 months of active surveillance followed by hospital-based surveillance. Thus, duration of protection against VCD cannot be assessed. However, active surveillance is currently being reinstated. Data on hospitalized dengue has been collected throughout the trial period, though with different surveillance systems in the Active and Hospital Phases. With the limitations of this change in surveillance and that the CYD and placebo groups have different histories of dengue exposure at the start of later time intervals, it is one source of data available now to assess protection over the period of the trial.

The methods used for re-analysis of the Phase 3 trial data are based on assays and statistical methods that are associated with misclassification of serostatus at baseline, which vary by assay. The false-positive rate (misclassifying seronegatives as serpositives) is high (31%), which would bias the result towards the null. Thus the protective effect in seropositives may be greater than that estimated in these analyses.