We report emerging subtropical bimodal seasonality and alternating predominance of norovirus GII.4 and non-GII.4 genotypes in Hong Kong. GII.4 predominated in summer and autumn months and affected young children, whereas emergent non-GII.4 genotypes predominated in winter months and affected all age groups. This highly dynamic epidemiology should inform vaccination strategies.

**The Study**

Hong Kong is a subtropical coastal city in southern China and has a temperate climate with hot summers and dry winters (Köppen-Geiger climate classification “Cwa”). Since March 2014, we have been monitoring virus genotypes in all hospitalized (i.e., severe) norovirus AGE case-patients of all ages diagnosed at the Prince of Wales Hospital of the Chinese University of Hong Kong and have reported the emergence and predominance of 2 previously less common non-GII.4 genotypes, GII.17 in 2014 and GII.2 in 2016 (8,9). Here we present further analysis of the seasonal dynamics of different norovirus genotypes during a 42-month period from March 2014 through August 2017. We identified norovirus genotypes in 1,100 (88.3%) of 1,246 case-patients by means of partial viral protein 1 Sanger sequencing and genotype assignment using a genotyping tool available through the National Institute of Public Health and the Environment of the Netherlands (http://www.rivm.nl/mpf/typingtool/norovirus). Seven case-patients were co-infected with >1 norovirus genotype. The proportion of GII.4 genotypes was 49.8% and that of non-GII.4 genotypes 50.2%. Overwhelmingly, most norovirus GII.4 belonged to the GII.Pe-GII.4 Sydney variant (512/544; 94.1%; online Technical Appendix Figure 1, https://wwwnc.cdc.gov/EID/article/24/4/17-1791-Techapp1.pdf). The recent recombinant GII.P16-GII.4 Sydney that emerged and predominated in the United States during 2016–2017 (10) was only detected sporadically in Hong Kong (online Technical Appendix Figure 1). The 2 most prevalent norovirus non-GII.4 genotypes were the recently emerged GII.17 (35.9%) and GII.2 (26.0%) (online Technical Appendix Figure 1).

We observed a bimodal seasonality of norovirus AGE requiring hospitalization, with periodic oscillation in the age group of admitted case-patients (Figure, panel A). Among the 19 months that norovirus preferentially affected young children <5 years old (as indicated by a monthly median age of case-patients <5 years), 17 (89%) were predominated by GII.4 genotype, whereas among the 23 months that preferentially affected older children and adults, 19 (83%) were predominated by non-GII.4 genotypes (Figure, panel B). By age groups, norovirus GII.4 accounted for most (68.5%) case-patients who were young children <5 years old, whereas norovirus non-GII.4 predominated in all other age groups: 5–17 years (75.7%), 18–40 years (87.0%), 41–65 years (78.6%), and >65 years (63.2%). The median age of case-patients infected with the recently emerged GII.17 and GII.2 was significantly higher than that for those infected with GII.4.
(GII.4, 2 years [interquartile range (IQR) 1–4 years]; GII.17, 49 years [IQR 10–72 years]; GII.2, 5 years [IQR 2–22 years]; p<0.0001, Kruskal-Wallis test) (online Technical Appendix Figure 2), as reported earlier over a shorter period (8,9). By season, late summer and autumn peaks were associated with norovirus GI.4, whereas winter peaks were associated with norovirus non-GI.4 (Figure, panel B). Norovirus infections have become equally common during summer and autumn months (52.3% of all infections during June–November) and during winter and spring months (47.7% of all infections during December–May).

Conclusions
We observed an influenza-like subtropical bimodal seasonality and alternating predominance of norovirus GI.4 and non-GI.4 genotypes, with each infecting different age groups. Norovirus GI.4 predominated in summer and autumn months and preferentially affected young children, who are also one of the age groups most affected by seasonal influenza. In contrast, emergent norovirus non-GI.4 predominated in winter months and affected wider age groups (e.g., all age groups were affected by GI.17 and older children and young adults by GI.2), a pattern which is reminiscent of pandemic influenza viruses. These findings illustrate a highly dynamic epidemiology of norovirus. A similar pattern of alternating epidemics has been observed among the 4 dengue virus serotypes and was shown to reflect moderate but not weak or strong interserotypic cross-protective immunity (10). The alternating predominance of norovirus GI.4 and non-GI.4 genotypes in severe infections might reflect an equally complex virus–human immunologic interaction on individual and population levels. This might be explained at least in part by the recently proposed concept that groups norovirus genotypes into so-called immunotypes (12).

The out-of-phase oscillation in the demographic characteristics of norovirus patients admitted to our hospitals

Figure. Bimodal seasonality and alternating predominance of norovirus GI.4 and non-GI.4 genotypes in Hong Kong, China, 2014–2017. A) Temporal distribution of ages of patients hospitalized for norovirus gastroenteritis. Each dot represents 1 patient. Red horizontal bars indicate medians. B) Epidemic curve during the study period. All cases shown are stratified by norovirus viral protein 1 genotype. Pink shading along baseline indicates months during which the median age of hospitalized case-patients was ≥5 years.
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Bimodal Seasonality and Alternating Predominance of Norovirus GII.4 and Non-GII.4, Hong Kong, China, 2014–2017

Technical Appendix

Technical Appendix Figure 1. Epidemic curve showing bimodal seasonality and alternating predominance of norovirus GII.4 and non-GII.4 (stratified by viral protein 1 genotype) in Hong Kong, China, 2014–2017. Pink shading along baseline indicates months during which the median age of hospitalized case-patients was >5 years as in Figure.
Technical Appendix Figure 2. Box-plot of age distribution of case-patients infected with norovirus viral protein 1 genotypes GII.4, GII.17, and GII.2. ****, two-tailed p<0.0001, as calculated by nonparametric Kruskal-Wallis test and corrected for Dunn’s multiple comparisons test.