

CORONAVIRUS

The Huanan Seafood Wholesale Market in Wuhan was the early epicenter of the COVID-19 pandemic

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Understanding how severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) emerged in 2019 is critical to preventing future zoonotic outbreaks before they become the next pandemic. The Huanan Seafood Wholesale Market in Wuhan, China, was identified as a likely source of cases in early reports, but later this conclusion became controversial. We show here that the earliest known COVID-19 cases from December 2019, including those without reported direct links, were geographically centered on this market. We report that live SARS-CoV-2-susceptible mammals were sold at the market in late 2019 and that within the market, SARS-CoV-2-positive environmental samples were spatially associated with vendors selling live mammals. Although there is insufficient evidence to define upstream events, and exact circumstances remain obscure, our analyses indicate that the emergence of SARS-CoV-2 occurred through the live wildlife trade in China and show that the Huanan market was the epicenter of the COVID-19 pandemic.

On 31 December 2019, the Chinese government notified the World Health Organization (WHO) of an outbreak of severe pneumonia of unknown etiology in Wuhan, Hubei Province, China (1–4), a city of ~11 million people. Of the initial 41 people hospitalized with unknown pneumonia by 2 January 2020, 27 (66%) had direct exposure to the Huanan Wholesale Seafood Market (hereafter, “Huanan market”) (2, 5, 6). These first cases were confirmed to be infected with a novel coronavirus, subsequently named severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), and were suffering from a disease later named coronavirus disease 2019 (COVID-19). The initial diagnoses of COVID-19 were made in several hospitals independently between 18 and 29 December 2019 (5). These early reports were free from ascertainment bias because they were based on signs and symptoms before the Huanan market was identified as a shared risk factor (5). A subsequent systematic review of all cases reported to China’s National Notifiable Disease Reporting System by hospitals in Wuhan

as part of the joint WHO-Chinese “WHO-convened global study of origins of SARS-CoV-2: China Part” (hereafter, “WHO mission report”) (7) showed that 55 of 168 of the earliest known COVID-19 cases were associated with this market. However, the observation that the preponderance of early cases were linked to the Huanan market, alone, does not establish that the pandemic originated there.

Sustained live mammal sales during 2019 occurred at the Huanan market and three other markets in Wuhan, and included wild and farmed wildlife (8). Several of these species are known to be experimentally susceptible to SARS-related coronaviruses (SARSR-CoVs) such as SARS-CoV (hereafter, “SARS-CoV-1”) and SARS-CoV-2 (9–11). During the early stages of the COVID-19 pandemic, animals sold at the Huanan market were hypothesized to be the source of the unexplained pneumonia cases (12–19) (data S1), consistent with the emergence of SARS-CoV-1 from 2002 to 2004 (20), as well as other viral zoonoses (21–23). This led to the decision to close and sanitize the Huanan market on 1 January 2020, with environmental

samples also being collected from vendors’ stalls (7, 12, 24) (data S1).

Determining the epicenter of the COVID-19 pandemic at the neighborhood level rather than at the city level could help to resolve whether SARS-CoV-2 had a zoonotic origin, similar to SARS-CoV-1 (20). In this study, we obtained data from a range of sources to test the hypothesis that the COVID-19 pandemic began at the Huanan market. Despite limited testing of live wildlife sold at the market, collectively, our results provide evidence that the Huanan market was the early epicenter of the COVID-19 pandemic and suggest that SARS-CoV-2 likely emerged from the live wildlife trade in China. However, events upstream of the market, as well as exact circumstances at the market, remain obscure, highlighting the need for further studies to understand and lower the risk of future pandemics.

Results

Early cases lived near to and centered on the Huanan market

The 2021 WHO mission report identified 174 COVID-19 cases in Hubei Province in December 2019 after careful examination of reported case histories (7). Although geographical coordinates of the residential locations of the 164 cases who lived within Wuhan were unavailable, we were able to reliably extract the latitude and longitude coordinates of 155 cases from maps in the report (figs. S1 to S8).

Although early COVID-19 cases occurred across Wuhan, most clustered in central Wuhan near the west bank of the Yangtze River, with a high density of cases near to, and surrounding, the Huanan market (Fig. 1A). We used a kernel density estimate (KDE) to reconstruct an underlying probability density function from which the home locations for each case were drawn (25). Using all 155 of the December 2019 cases, the location of the Huanan market lies within the highest density contour that contains 1% of the probability mass (Fig. 1B). For a KDE estimated using the 120 cases with no known linkage to the market, the market remains within the highest density 1% contour (Fig. 1C). The clustering of COVID-19 cases in December around the Huanan market (Fig. 1, B and C,

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insets) contrasts with the pattern of widely dispersed cases across Wuhan by early January through mid-February 2020 (Fig. 1, D and E), which we mapped using location data from individuals who had used a COVID-19 assistance channel on Sina Weibo, a Chinese social media platform (26). Weibo-based data analyses showed that, unlike early COVID-19 cases, by January and February, many of the sick individuals who sought help resided in highly populated areas of the city, particularly in areas with a high density of older people (Fig. 1E and figs. S9 and S10).

We also investigated whether the December COVID-19 cases were closer to the market than expected based on an empirical null distribution of Wuhan's population density [data from WorldPop.org (27, 28)], with a median distance to the Huanan market of 16.11 km (25). To account for older individuals being more likely to be hospitalized and sick with COVID-19 (29), we age-matched the population data to the December 2019 COVID-19 case data. We considered three categories of cases, which were all significantly closer to the Huanan market than expected: (i) all cases (median distance 4.28 km; $P < 0.001$), (ii) cases linked directly to the Huanan market (median distance 5.74 km; $P < 0.001$), and (iii) cases with no evidence of a direct link to the Huanan market (median distance 4.00 km; $P < 0.001$) (Fig. 2A). The cases with no known link to the market on average resided closer to the market than the cases with links to the market ($P = 0.029$). Furthermore, the distances between the center points (Fig. 2B) and the Huanan market were shorter than expected for all categories of December cases compared with the empirical null distribution of Wuhan's population density (Fig. 2A). For all December cases, the center point was located 1.02 km away ($P = 0.007$); for cases with market links, it was 2.28 km away ($P = 0.034$); and for the cases with no reported link to the market, it was 0.91 km away ($P = 0.006$). By comparison, the center point of age-matched samples drawn from the empirical null distribution was 4.65 km away from the market (Fig. 2A).

We tested the robustness of our results for the possibility of ascertainment bias (25). For all mapped cases ($n = 155$), under the "center-point distance to the Huanan market" test, the 38 cases residing closest to the market (within a radius of 1.6 km) could be removed from the dataset before losing significance at the $\alpha = 0.05$ level (fig. S12). For the "median distance to Huanan market" test, we could remove 98 cases (63%) ($r = 5.8$ km). For cases not directly linked to the Huanan market ($n = 120$), we could remove 36 (30%) ($r = 1.5$ km) and 81 (68%) ($r = 4.3$ km) cases for the two tests, respectively, before losing significance at the $\alpha = 0.05$ level (fig. S12).

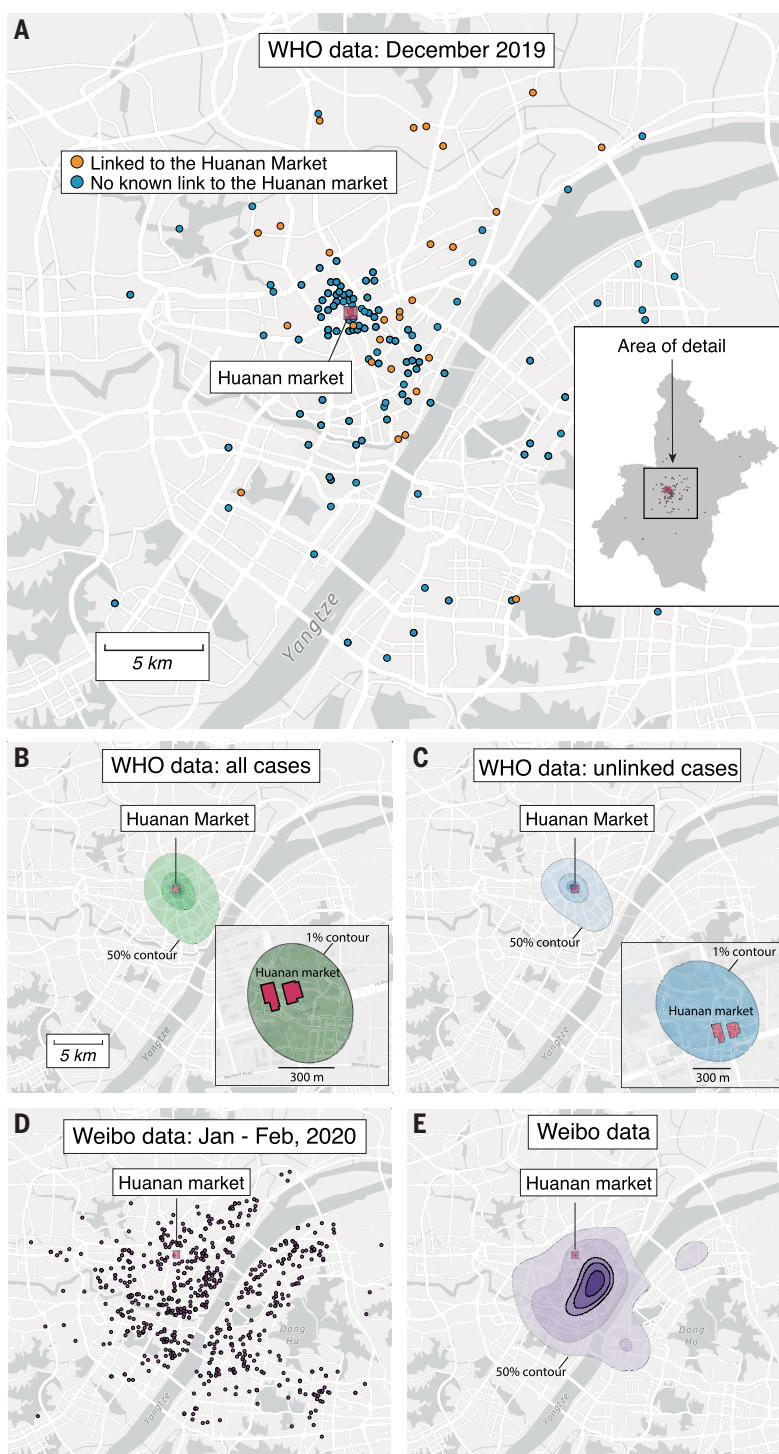
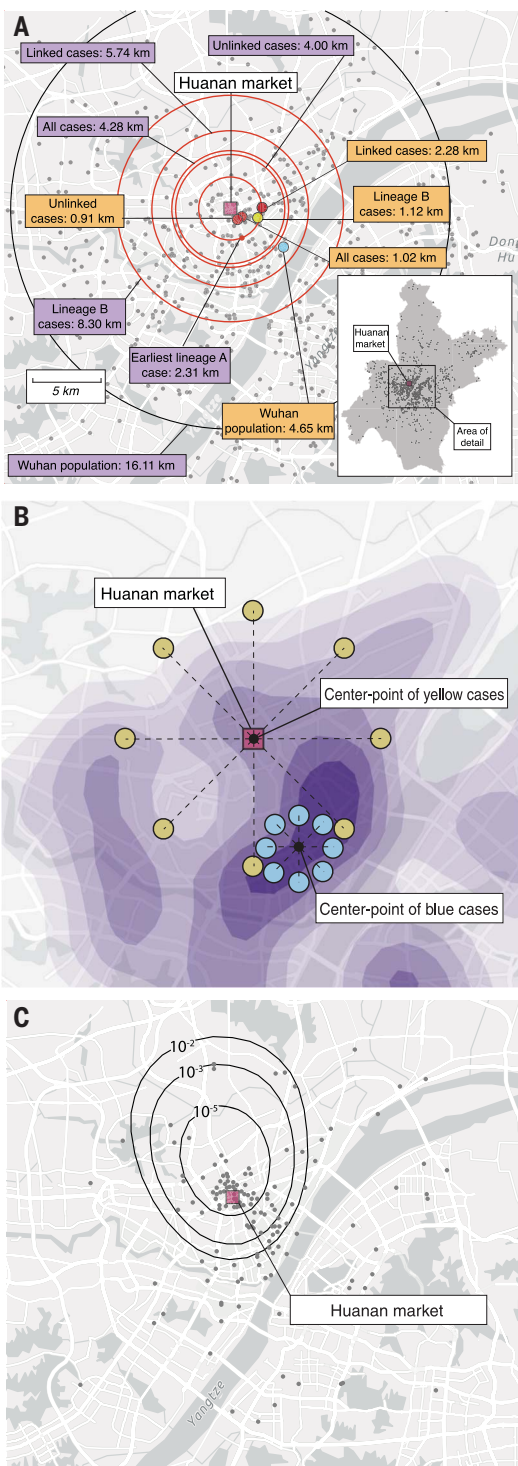


Fig. 1. Spatial patterns of COVID-19 cases in Wuhan in December 2019 and January–February 2020. (A) Locations of the 155 cases that we extracted from the WHO mission report (7). Inset: map of Wuhan with the December 2019 cases indicated with gray dots (no cases are obscured by the inset). In both the inset and the main panel, the location of the Huanan market is indicated with a red square. (B) Probability density contours reconstructed by a KDE using all 155 COVID-19 cases locations from December 2019. The highest density 50% contour marked is the area for which cases drawn from the probability distribution are as likely to lie inside as outside. Also shown are the highest density 25%, 10%, 5%, and 1% contours. Inset: expanded view and the highest density 1% probability density contour. (C) Probability density contours reconstructed using the 120 COVID-19 cases locations from December 2019 that were unlinked to the Huanan market. (D) Locations of 737 COVID-19 cases from Weibo data dating to January–February 2020. (E) The same highest probability density contours (50% through 1%) as shown in (B) and (C) for 737 COVID-19 case locations from Weibo data.

Fig. 2. Spatial analyses. (A) Inset: map of Wuhan, with gray dots indicating the 1000 random samples from the WorldPop.com null distribution. In the main panel, the median distance between Huanan market and the WorldPop.org null distribution is indicated by the outer black circle. December 2019 cases are indicated by concentric red circles (distances to Huanan market are described in the purple boxes). The center point of Wuhan population density data is indicated by a blue dot. Center points of December 2019 case locations are shown as follows: red dots indicate “all,” “linked,” and “unlinked” cases, and the yellow dot indicates lineage B cases. Distance from center points to Huanan market are described in orange boxes. (B) Schematic showing how cases can be near to, but not centered on, a specific location. We hypothesized that if the Huanan market were the epicenter of the pandemic, then early cases should fall not just unexpectedly near to it but should also be unexpectedly centered on it (see the materials and methods). The blue dots show how hypothetical cases quite near the Huanan market could nevertheless not be centered on it. (C) Tolerance contours based on relative risk of COVID-19 cases in December 2019 versus data from January–February 2020. The gray dots show the December case locations. The contours represent the probability of observing that density of December cases within the bounds of the given contour if the December cases had been drawn from the same spatial distribution as the January–February data.



We performed a spatial relative risk analysis (25) to compare December 2019 COVID-19 cases with January–February 2020 cases reported through Weibo (Fig. 2C). The Huanan market is located within a well-defined area with high case density that would be expected to be observed in <1 in 100,000 samplings of the Weibo data empirical distribution (the relative risk analysis is shown in Fig. 2C and the control

distribution in Fig. 1D). No other regions in Wuhan showed a comparable case density.

Both early lineages of SARS-CoV-2 were geographically associated with the market

Two lineages of SARS-CoV-2 designated A and B (30) have co-circulated globally since early in the COVID-19 pandemic (31). Until a report in a recent preprint (24), only lineage B sequen-

ces had been sampled at the Huanan market. The 11 lineage B cases from December 2019 for which we have location information resided closer than expected to the Huanan market compared with the age-matched Wuhan population distribution (median distance 8.30 km; $P = 0.017$) (25). The center point of the 11 lineage B cases was 1.95 km from the Huanan market, also closer than expected ($P = 0.026$). The two lineage A cases for which we have location information involved the two earliest lineage A genomes known to date. Neither case reported any contact with the Huanan market (7). The first case was detected before any knowledge of a possible association of the unexplained pneumonia in Wuhan with the Huanan market (5), and therefore could not have been a product of ascertainment bias in favor of cases residing near the market. The second case had stayed in a hotel near the market (32) for the 5 days preceding symptom onset (25). Relative to the age-matched Wuhan population distribution, the first individual resided closer to the Huanan market (2.31 km) than expected ($P = 0.034$). Although the exact location of the hotel near the market was not reported (32), there are at least 20 hotels within 500 m (table S1). Under the conservative assumption that the hotel could have been located as far as 2.31 km from the Huanan market (as was the residence of the other lineage A case), and assuming that this location is comparable to a residential location given the timing of the stay before symptom onset (25), it would be unlikely to observe both of the earliest lineage A cases this near to the Huanan market ($P = 0.001$ or less). The finding that both identified lineage A cases had a geographical connection to the market, in combination with the detection of lineage A within the market (24), support the likelihood that during the early epidemic, lineage A was, like lineage B, disseminating outward from the Huanan market into the surrounding neighborhoods.

Our statistical results were robust to a range of factors, for example, the use of an empirical control distribution that was based on presumptive COVID-19 cases locations later in the Wuhan epidemic (Weibo data); laboratory-confirmed versus clinically diagnosed cases; and uncertainty in case location or missing data (figs. S13 to S15) (25). For instance, we artificially introduced location uncertainty (“noise”) in each case location in our dataset by randomly resampling each point within a circle of radius 1000 m centered on its original center point, and the conclusions were unaffected (fig. S13). The extraction method that we used actually introduced only up to ~50 m of noise in each case location estimate (fig. S7), ruling out the possibility that our overall results were affected by this source of error. The results were also robust when corrected for multiple-hypothesis testing (table S4).

Wild animal trading in Wuhan markets

In addition to selling seafood, poultry, and other commodities, the Huanan market was among four markets in Wuhan reported to consistently sell a variety of live wild-captured or farmed mammal species in the years and months leading up to the COVID-19 pandemic (8). There are, however, no prior reports of which species, if any, were sold at the Huanan market in the months leading up to the pandemic. Here, we report that multiple plausible intermediate wildlife hosts of SARS-CoV-2 progenitor viruses, including red foxes (*Vulpes vulpes*), hog badgers (*Arctonyx albogularis*), and common raccoon dogs (*Nyctereutes procyonoides*), were sold live at the Huanan market up until at least November 2019 (Table 1 and table S5). No reports are known to be available for SARS-CoV-2 test results from these mammals at the Huanan market. Despite a general slowdown in live animal sales during the winter months, we report that raccoon dogs, which are sold for both meat and fur, were consistently available for sale throughout the year, including at the Huanan market in November 2019 (Table 1 and table S5).

There were potentially many locations in Wuhan, a city of 11 million, that would have been equally or more likely than the Huanan market to sustain the first recognized cluster of a new respiratory pathogen had its introduction not been linked to a live animal market, including other shopping venues, hospitals, elder care facilities, workplaces, universities, and places of worship. To investigate possi-

ble sites, we compared the relative extent of intra-urban human traffic to the Huanan market versus other locations within the city of Wuhan using a location-specific dataset of social media check-ins in the Sina Visitor System (25, 33). We found at least 70 other markets throughout the city of Wuhan that received more social media check-ins than the Huanan market (Fig. 3). To extend this analysis beyond only markets, we also used a subsequently published list of known SARS-CoV-2 superspread locations (34) to identify 430 locations in Wuhan that may have been at high risk for superspread events and which received more check-ins than the Huanan market (Fig. 3, inset). The Huanan market accounted for 0.12% (120 of 98,146) of social media check-ins to markets in the dataset that received at least as many check-ins as the Huanan market. The market accounted for 0.04% (120 of 262,233) of all social media check-ins to the >400 sites in Wuhan identified as especially likely to be potential superspreader locations and which received at least as many social media visits as the Huanan market. Considering the number of check-ins to all four markets selling live wild animals in Wuhan (combined), they accounted for 0.21% (206 of 98,146) of market visits and 0.079% (206 of 262,233) of visits to the 430 potential superspread sites, where a new respiratory disease might first be noticed in a large city.

A dataset from the Chinese Center for Disease Prevention and Control (CCDC) report dated 22 January 2020 (data S1) (12, 13, 15, 16) was

made publicly available in June 2020 (24, 35). A total of 585 environmental samples were initially taken from various surfaces in the Huanan market on 1 and 12 January 2020 by the CCDC (tables S6 and S7 and data S1) (12, 13, 15, 16, 24, 35), with further samples taken throughout the market during January and February (24). We extended the analysis in the WHO mission report (7) by integrating public online maps and photographic evidence, data from public business registries (table S8 and data S2), information about which live mammal species were sold at the Huanan market in late 2019 (Table 1 and table S5), and the CCDC report (data S1). We reconstructed the floor plan of the market and integrated information from business registries of vendors at the market (fig. S16 and table S8), as well as an official report (36) recording fines to three business owners for illegal sales of live mammals (data S2) (36). From this, we identified an additional five stalls that were likely selling live or freshly butchered mammals or other unspecified meat products in the southwest corner of the western section of the market (Fig. 4A, figs. S16 and S17, and table S6).

Five of the SARS-CoV-2-positive environmental samples were taken from a single stall selling live mammals in late 2019 (table S6). Further, all five objects sampled showed an association with animal sales, including a metal cage, two carts (of the kind frequently used to transport mobile animal cages), and a hair and feather remover (table S6). No human COVID-19 cases were reported there (7, 12). The same

Table 1. Live mammals traded at the Huanan market in November and December 2019.

Species (susceptibility*)	Family (susceptibility*)	Order (susceptibility*)	Observed at Huanan market November 2019
Raccoon dog (<i>Nyctereutes procyonoides</i>) (Y)	Canidae (Y)	Carnivora (Y)	Y
Amur hedgehog (<i>Erinaceus amurensis</i>)	Erinaceidae	Eulipotyphla	Y
Hog badger (<i>Arctonyx albogularis</i>) (Y)	Mustelidae (Y)	Carnivora (Y)	Y
Asian badger (<i>Meles leucurus</i>)	Mustelidae (Y)	Carnivora (Y)	Y
Chinese hare (<i>Lepus sinensis</i>)	Leporidae (Y)	Lagomorpha (Y)	Y
Chinese bamboo rat (<i>Rhizomys sinensis</i>) (Y)	Spalacidae (Y)	Rodentia (Y)	Y
Malayan porcupine (<i>Hystrix brachyura</i>)	Hystricidae	Rodentia (Y)	Y
Chinese muntjac (<i>Muntiacus reevesi</i>)	Cervidae (Y)	Artiodactyla (Y)	Y
Marmot (<i>Marmota himalayana</i>)	Sciuridae	Rodentia (Y)	Y
Red fox (<i>Vulpes vulpes</i>) (Y)	Canidae (Y)	Carnivora (Y)	Y
Siberian weasel (<i>Mustela sibirica</i>)	Mustelidae (Y)	Carnivora (Y)	N†
Pallas's squirrel (<i>Callosciurus erythraeus</i>)	Sciuridae	Rodentia (Y)	N
Masked palm civet (<i>Paguma larvata</i>) (Y)	Viverridae (Y)	Carnivora (Y)	N
Coypu (<i>Myocastor coypus</i>)	Echimyidae	Rodentia (Y)	N
Mink (<i>Neovison vison</i>) (Y)	Mustelidae (Y)	Carnivora (Y)	N
Red squirrel (<i>Sciurus vulgaris</i>)	Sciuridae	Rodentia (Y)	N
Wild boar (<i>Sus scrofa</i>) (Y)	Suidae (Y)	Artiodactyla (Y)	N
Complex-toothed flying squirrel (<i>Trogopterus xanthipes</i>)	Sciuridae	Rodentia (Y)	N

*Based on live susceptibility findings, serological findings, or ACE2-binding assays. See table S5 for details and associated references. Wuhan market during the 2017–2019 study period (8).

†Animals listed as "N" (no) were, however, present at

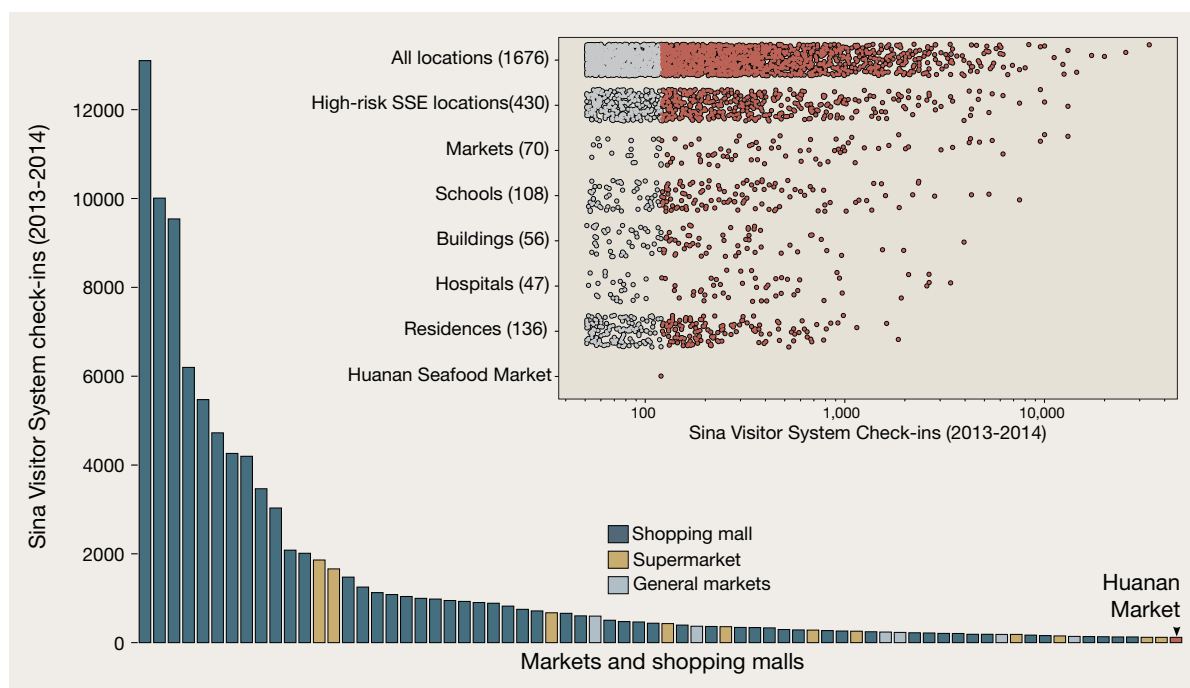


Fig. 3. Visitors to locations throughout Wuhan. Shown is the number of social media check-ins in the Sina Visitor System from 2013 to 2014 as shared by (33). The numbers of check-ins to individual markets throughout the city are shown in comparison with check-ins at the Huanan market. Inset: the total

number of check-ins to all individual locations across the city of Wuhan grouped by category. Locations with >50 visitor check-ins are shown, and the locations that received more check-ins than the Huanan market in the same period are shown in red.

stall was visited by one of us (E.C.H.) in 2014, and live raccoon dogs were observed housed in a metal cage stacked on top of a cage with live birds (Fig. 4A) (37). A recent report (24) identified that the grates outside of this stall, upon which animal cages were stacked (37), were positive for SARS-CoV-2.

Positive environmental samples linked both to live mammal sales and to human cases at the Huanan market

We used a spatial relative risk analysis to identify potential regions of the market with an increased density of positive environmental samples (25). We found evidence ($P < 0.05$) of a region in the southwest area of the market where live mammals were for sale (Fig. 4B). Although environmental sampling of the market was incomplete and spatially heterogeneous (data S1 and table S6), our analysis accounts for the empirical environmental sampling distribution, which was biased toward “stalls related to December cases,” as well as “stalls that sold livestock, poultry, farmed wildlife” (7) (Fig. 4, C and D). The “distance to the nearest vendor selling live mammals” and the “distance to the nearest human case” were independently predictive of environmental sample positivity ($P = 0.004$ and 0.014 , respectively, for $n = 6$; table S9). To further investigate the robustness of these findings to possible sampling biases, we considered

three scenarios: (i) oversampling of live mammal and unknown meat stalls, (ii) overcounting of positive samples, and (iii) exclusion of the seafood stand near the wildlife area of the market (with five positive samples) from our analysis (table S10). In each case, the distance to live mammal vendors remained predictive of environmental sample positivity, and the region of increased positive sample density in the southwest corner of the western section of the market remained consistent (fig. S18).

Finally, to analyze the spatial patterning of human cases within the Huanan market, we plotted cases as a function of symptom onset from the WHO mission report (7) (Fig. 5A and table S11) (25). All eight COVID-19 cases detected before 20 December 2019 were from the western side of the market, where mammal species were also sold (Fig. 5, B and C). Unlike SARS-CoV-2-positive environmental samples (Fig. 4, A and C), we found that COVID-19 cases were more diffuse throughout the building (Fig. 5).

Study limitations

There are several limitations to our study. We have been able to recover location data for most of the December-onset COVID-19 cases identified by the WHO mission (7) with sufficient precision to support our conclusions. However, we do not have access to the precise latitude and longitude coordinates of all of these cases. Should such data exist, they may

be accompanied by additional metadata, some of which we have reconstructed, but some of which, including the date of onset of each case, would be valuable for ongoing studies. We also lack direct evidence of an intermediate animal infected with a SARS-CoV-2 progenitor virus either at the Huanan market or at a location connected to its supply chain, such as a farm. Additionally, no line list of early COVID-19 cases is available, and we do not have complete details of environmental sampling. However, compared with many other outbreaks, we have more comprehensive information on early cases, hospitalizations, and environmental sampling (7).

Discussion

Several lines of evidence support the hypothesis that the Huanan market was the epicenter of the COVID-19 pandemic and that SARS-CoV-2 emerged from activities associated with the live wildlife trade there. Spatial analyses within the market show that SARS-CoV-2-positive environmental samples, including cages, carts, and freezers, were associated with activities concentrated in the southwest corner of the market. This is the same section where vendors were selling live mammals, including raccoon dogs, hog badgers, and red foxes, immediately before the COVID-19 pandemic. Multiple positive samples were taken from one stall known to have sold live

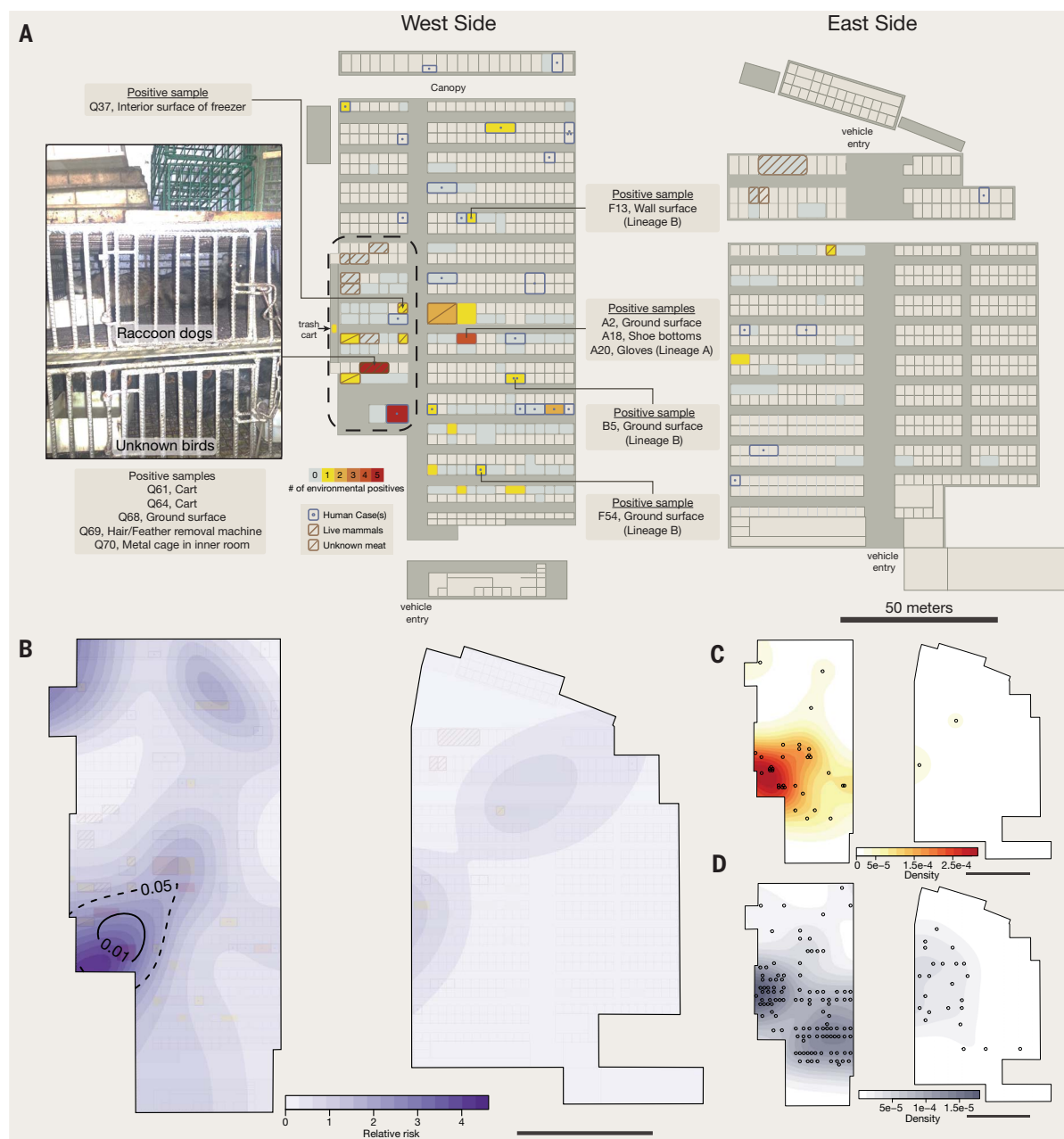


Fig. 4. Map of the Huanan market. (A) Aggregated environmental sampling and human case data from the Huanan market. Captions describe the types of SARS-CoV-2-positive environmental samples obtained from known live animal vendors (left) and from stalls with samples with known virus lineage (center). Lineage is unknown unless noted; sequencing data have not been released for some samples, and many samples were PCR-positive but not sequenced. Image at left shows raccoon dogs in a metal cage on top of caged birds from a business with five positive environmental samples (photo by E.C.H.). Center: Rectangle with dashed outline indicates the “wildlife” section of the market.

(B) Relative risk analysis of positive environmental samples. Tolerance contours enclose regions with statistically significant elevation in density of positive environmental samples relative to the distribution of sampled stalls. (C) Distribution of positive environmental samples. Sample locations (centroid of corresponding business) and quantity are shown as black circles. (D) Control distribution for relative risk analysis. All businesses investigated with environmental sampling are shown as black circles (there is one circle per business regardless of whether a positive sample was found). See table S12 for details on stalls that were SARS-CoV-2-negative.

mammals, and the water drain proximal to this stall, as well as other sewerages and a nearby wildlife stall on the southwest side of the market, tested positive for SARS-CoV-2 (24). These findings suggest that infected animals were present at the Huanan market at the

beginning of the COVID-19 pandemic; however, we do not have access to any live animal samples from relevant species. Additional information, including sequencing data and detailed sampling strategy, would be invaluable to test this hypothesis comprehensively.

In a related study, we inferred separate introductions of SARS-CoV-2 lineages A and B into humans from likely infected animals at the Huanan market (38). We estimated the first COVID-19 case to have occurred in November 2019, with few human cases and hospitalizations

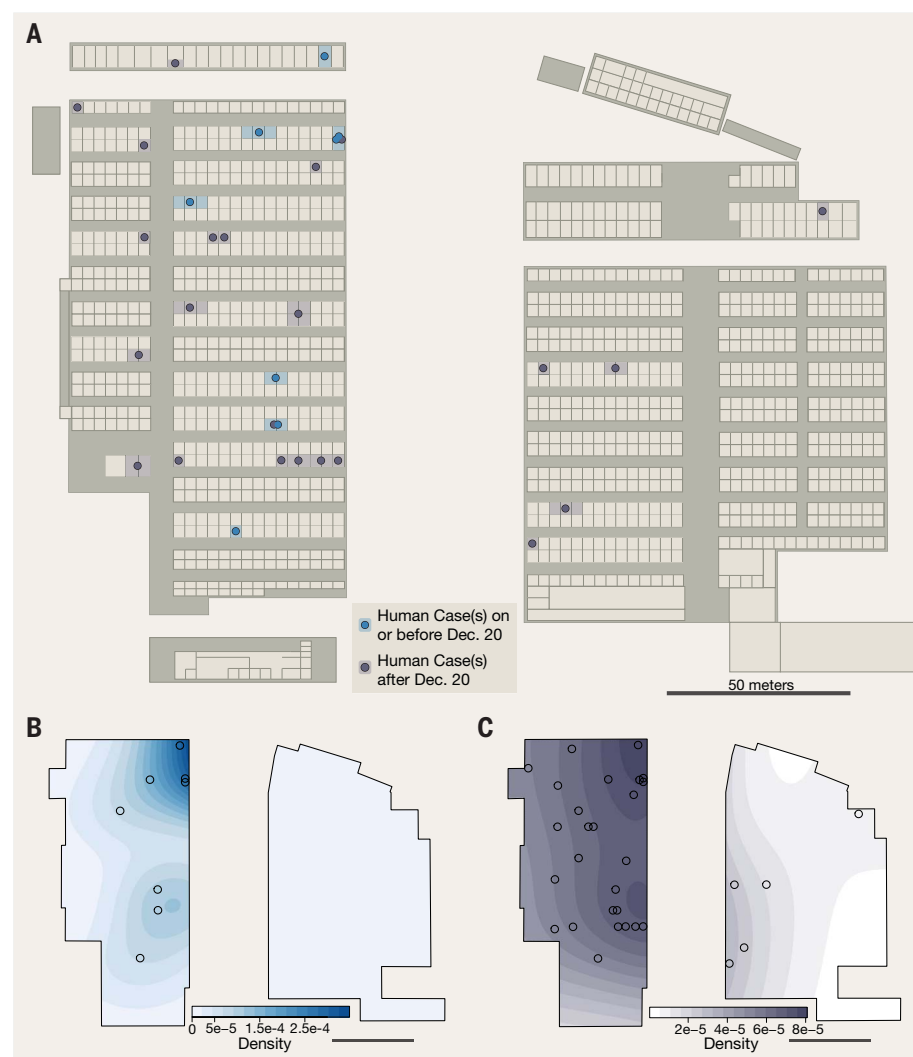


Fig. 5. Location and timing of human cases in Huanan market. (A) Outline colors correspond to the timing of the first known case in each business. Individual case timing is denoted by marker color and shown within the outlined business. (B) Distribution of known cases on or before 20 December 2019. Case locations are shown as black circles. (C) Distribution of all known human cases in Huanan market. See table S11 for details on SARS-CoV-2-positive human cases with the Huanan market.

occurring through mid-December (38). A recent preprint (24) confirms the authenticity of the CCDC report (data S1) and records additional positive environmental samples in the southwestern area of the market selling live animals. This report also documents the early presence of the A lineage of SARS-CoV-2 in a Huanan market environmental sample. This, along with the lineage A cases that we report in close geographical proximity to the market in December 2019, challenges the suggestion that the market was simply a superspreading event, which would be lineage specific. Rather, it adds to the evidence presented here that lineage A, like lineage B, may have originated at the Huanan market and then spread from this epicenter into the neighborhoods surrounding the market and beyond.

Several observations suggest that the geographic association of early COVID-19 cases with the Huanan market is unlikely to have been the result of ascertainment bias (see the supplementary text and tables S2 and S3) (39). These include that (i) few, if any, cases among Huanan market–unlinked individuals are likely to have been detected by active searching in the neighborhoods around the market, only in hospitals, because all of the cases analyzed here were hospitalized (7); (ii) public health officials simultaneously became aware of Huanan-linked cases both near and far from the Huanan market, not just the ones near it (fig. S11) (5); (iii) Huanan market–unlinked cases would not be expected to live significantly closer to the market than linked cases if they had been ascertained as contacts traced from those market-linked cases; and (iv) seroprevalence

in Wuhan was highest in the districts around the market (40, 41). It is also noteworthy that the December 2019 COVID-19 cases that we consider here were identified based on reviews of clinical signs and symptoms, not epidemiological factors such as where they resided or links to the Huanan market (7), and that excess deaths from pneumonia rose first in the districts surrounding the market (42). Moreover, the spatial relationship with the Huanan market remains after removing the two-thirds of the unlinked cases residing nearest the market.

One of the key findings of our study is that “unlinked” early COVID-19 patients, i.e., those who did not work at the market, did not know someone who did, and had not recently visited the market, resided significantly closer to the market than patients with a direct link to it. The observation that a substantial proportion of early cases had no known epidemiological link had previously been used as an argument against the Huanan market being the epicenter of the pandemic. However, this group of cases resided significantly closer to the market than those who worked there, indicating that they had been exposed to the virus at or near the Huanan market. For market workers, the exposure risk was their place of work, not their residential locations, which were significantly farther afield than those cases not formally linked to the market.

Our spatial analyses show how patterns of COVID-19 cases shifted between late 2019, when the outbreak began (43), and early 2020, as the epidemic spread widely across Wuhan. COVID-19 cases in December 2019 were associated with the Huanan market in a manner unrelated to Wuhan population density or demographic patterns, unlike the wide spatial distribution of cases observed during later stages of the epidemic in January–February 2020. This observation fits with the evidence from other sources that SARS-CoV-2 was not widespread in Wuhan at the end of 2019. For example, no SARS-CoV-2-positive sera or influenza-like illness reports were recorded among more than 40,000 blood donor samples collected up to December 2019 (44, 45), and none of thousands of samples taken from patients with influenza-like illness at Wuhan hospitals in October to December 2019 tested for SARS-CoV-2 RNA was positive (7).

The sustained presence of a potential source of virus transmission into the human population in late 2019, plausibly from infected live mammals sold at the Huanan market, offers an explanation of our findings and the origins of SARS-CoV-2. The pattern of COVID-19 cases reported for the Huanan market, with the earliest cases in the same part of the market as the wildlife sales and evidence of at least two introductions (38), resembles the multiple cross-species transmissions of SARS-CoV-2 subsequently observed during the pandemic from

animals to humans on mink farms (46) and from infected hamsters to humans in the pet trade (47). There was an extensive network of wildlife farms in western Hubei Province, including hundreds of thousands of raccoon dogs on farms in Enshi Prefecture, which supplied the Huanan market (48). This region of Hubei contains extensive cave complexes housing *Rhinolophus* bats, which carry SARS-CoVs (49). SARS-CoV-1 was recovered from farmed masked palm civets (*Paguma larvata*) from Hubei in 2003 and 2004 (20). The animals on these farms (nearly 1 million) were rapidly released, sold, or killed in early 2020 (48), apparently without testing for SARS-CoV-2 (7). Live animals sold at the market (Table 1) were apparently not sampled either. By contrast, during the SARS-CoV-1 outbreaks, farms and markets remained open for more than a year after the first human cases occurred, allowing sampling of viruses from infected animals (20).

The live animal trade and live animal markets are a common theme in virus spillover events (21–23, 50), with markets such as the Huanan market selling live mammals being in the highest risk category (51). The events leading up to the COVID-19 pandemic mirror the SARS-CoV-1 outbreaks from 2002 to 2004, which were traced to infected animals in the Guangdong, Jiangxi, Henan, Hunan, and Hubei provinces in China (20). Maximum effort must now be applied to elucidate the upstream events that might have brought SARS-CoV-2 into the Huanan market, culminating in the COVID-19 pandemic. To reduce the risk of future pandemics, we must understand, and then limit, the routes and opportunities for virus spillover.

Methods summary

Ethics statement

This research was reviewed by the Human Subject Protection Program at the University of Arizona and the Institutional Review Board (IRB) at The Scripps Research Institute and determined to be exempt from IRB approval because it constitutes secondary research for which consent is not required.

Data sources

COVID-19 case data from December 2019 were obtained from the WHO mission report (7) and from our previous analyses (5). Location information was extracted and sensitivity analyses performed to confirm accuracy and assess potential ascertainment bias. Geotagged January–February 2020 data from Weibo COVID-19 help seekers was obtained from the authors (26). Population density data were obtained from WorldPop.org (27). Sequencing- or quantitative polymerase chain reaction (PCR)-based environmental sample SARS-CoV-2 positivity from the Huanan market was

obtained from a January 2020 CCDC report (data S1) (24).

Wildlife trading at the Huanan market

Animal sales from Wuhan wet markets immediately before the COVID-19 pandemic were previously reported (8), and in this study we report details about animals for sale at the Huanan market up until November 2019.

Spatial analyses of COVID-19 cases

Haversine distances to the Huanan market were calculated for each of the geolocated December 2019 cases. Center points and median distances from cases to the Huanan market were calculated separately for (i) all 155 cases, (ii) the 35 cases epidemiologically linked to the Huanan market, (iii) the 120 cases not epidemiologically linked to the market, (iv) the 11 lineage B cases, and (v) the earliest lineage A case. These distances were also calculated for the 737 Weibo help seekers from 8 January to 10 February 2020 (26). Empirical null distributions were generated from the population density data and the Weibo data. The population density-null distributions were age-matched to the December 2019 cases. KDEs were also generated for the market-linked cases, unlinked cases, and all cases to infer a probability density function from which the cases could have been drawn. Highest-density contours representing specific probability masses (0.5, 0.25, 0.1, 0.05, and 0.01) were inferred, and the location of the market was compared with these.

Mobility analyses

To estimate the relative amount of intra-urban human traffic to the Huanan market compared with other locations within the city of Wuhan, we used a location-specific dataset of social media check-ins in the Sina Visitor System as shared by Li *et al.* (33). This dataset is based on 1,491,499 individual check-in events across the city of Wuhan from the years 2013–2014 (5 to 6 years before the start of the COVID-19 pandemic), and 770,521 visits were associated with 312,190 unique user identifiers. Location names and categories were translated using a Python API for Google Translate.

Spatial analyses of environmental samples at the Huanan market

We used the official maps from the CCDC (J2) (data S1) and the WHO map (7), as well as satellite photographs (Google Maps, Google Earth, Baidu Maps), aerial photographs, and images of the market in the public domain to reconstruct the floorplan of the market. Market stalls were assigned by categories of the types of goods sold using official reports and data from the TianYanCha.com business directory (this company has since gone out of business; for screenshots, see table S8 and data S2). Final maps of the Huanan market were converted

into geoJSON format for spatial analyses. Significance testing of live animal vendors and/or human SARS-CoV-2 cases on the number of positive environmental samples was performed using a binomial general linear model. Distances between businesses were defined as the distance between their respective center points, and spatial relative risk analysis was performed using the ‘sparr’ package in R, with linear boundary kernels for edge correction (52) and bandwidth selection performed using least-squares cross-validation.

REFERENCES AND NOTES

1. Sina Finance, “Wuhan pneumonia of unknown cause cases isolated, test results to be announced ASAP” (Sina Finance, 2019); <https://finance.sina.cn/2019-12-31/detail-ijhnzhak1074832.d.html?from=wap>.
2. Wuhan Municipal Health Commission, “Wuhan Municipal Health Commission’s briefing on the current situation of pneumonia in our city” (Wuhan Municipal Health Commission, 2019); <https://web.archive.org/web/20200131202951/http://wjw.wuhan.gov.cn/front/web/showDetail/2019123108989>.
3. World Health Organization, “COVID-19 – China” (WHO, 2020); <https://www.who.int/emergencies/diseases-outbreak-news/item/2020-DON229>.
4. The Novel Coronavirus Pneumonia Emergency Response Epidemiology Team, *China CDC Wkly* **2**, 113–122 (2020).
5. M. Worobey, *Science* **374**, 1202–1204 (2021).
6. C. Huang *et al.*, *Lancet* **395**, 497–506 (2020).
7. World Health Organization, “WHO-convened global study of origins of SARS-CoV-2: China Part” (WHO, 2021); <https://www.who.int/publications/item/who-convened-global-study-of-origins-of-sars-cov-2-china-part>.
8. X. Xiao, C. Newman, C. D. Buesching, D. W. Macdonald, Z.-M. Zhou, *Sci. Rep.* **11**, 11898 (2021).
9. C. M. Freuling *et al.*, *Emerg. Infect. Dis.* **26**, 2982–2985 (2020).
10. W. K. Jo *et al.*, *Transbound. Emerg. Dis.* **68**, 1824–1834 (2021).
11. I. R. Fischhoff, A. A. Castellanos, J. P. G. L. M. Rodrigues, A. Varsani, B. A. Han, *Proc. Biol. Sci.* **288**, 20211651 (2021).
12. W. Guizhen, “Chinese CDC disease control report” (see data S1).
13. Xinhua News, “Good news! Phased progress made in tracing the origin of the coronavirus” (Xinhua News, 2020); http://www.xinhuanet.com/politics/2020-01/26/c_1125503792.htm.
14. Beijing News, “Huanan Seafood Market in the pneumonia of unexplained incident” (Beijing News, 2020); <http://www.bjnews.com.cn/feature/2020/01/02/669054.html>.
15. Chinese Center for Disease Control and Prevention, “Chinese Center for Disease Control and Prevention detects large quantity of novel coronavirus in Wuhan Huanan Seafood Market” (Chinese CDC, 2020); https://www.chinacdc.cn/yw_9324/202001/t20200127_211469.html.
16. Yicai Global, “China detects large quantity of novel coronavirus at Wuhan Seafood Market” (Yicai Global, 2020); <https://www.yicaiglobal.com/opinion/yicai.global/china-detects-large-quantity-of-novel-coronavirus-at-wuhan-seafood-market>.
17. Chinese Center for Disease Control and Prevention, “China CDC calls on the public to protect themselves” (Chinese CDC, 2020); https://www.chinacdc.cn/yw_9324/202001/t20200128_211498.html.
18. Chinese Center for Disease Control and Prevention, “On the front line, disease control warriors race against the new coronavirus” (Chinese CDC, 2020); https://www.chinacdc.cn/yw_9324/202002/t20200201_212137.html.
19. Xinhua News, “China detects large quantity of novel coronavirus at Wuhan seafood market” (Xinhua News, 2020); https://web.archive.org/web/20200126230041/http://www.xinhuanet.com/english/2020-01/27/c_138735677.htm.
20. Z. Shi, Z. Hu, *Virus Res.* **133**, 74–87 (2008).
21. W. B. Karesh, R. A. Cook, E. L. Bennett, J. Newcomb, *Emerg. Infect. Dis.* **11**, 1000–1002 (2005).
22. N. D. Wolfe, P. Daszak, A. M. Kilpatrick, D. S. Burke, *Emerg. Infect. Dis.* **11**, 1822–1827 (2005).
23. C. K. Johnson *et al.*, *Proc. Biol. Sci.* **287**, 20192736 (2020).

24. G. Gao *et al.*, "Surveillance of SARS-CoV-2 in the environment and animal samples of the Huanan Seafood Market" [Preprint] (Research Square, 2022); <https://www.researchsquare.com/article/rs-1370392/v1>.
25. Material and methods are available as supplementary materials.
26. Z. Peng, R. Wang, L. Liu, H. Wu, *ISPRS Int. J. Geoinf.* **9**, 402 (2020).
27. WorldPop, "WorldPop: Open spatial demographic data and research" (2020); <http://WorldPop.org>.
28. A. J. Tatem, *Sci. Data* **4**, 170004 (2017).
29. M. O'Driscoll *et al.*, *Nature* **590**, 140–145 (2021).
30. A. Rambaut *et al.*, *Nat. Microbiol.* **5**, 1403–1407 (2020).
31. outbreak.info, "SARS-CoV-2 (nCoV-19) mutation reports: Lineage/mutation tracker" (outbreak.info, 2022); <https://outbreak.info/situation-reports>.
32. R. Lu *et al.*, *Lancet* **395**, 565–574 (2020).
33. L. Li, L. Yang, H. Zhu, R. Dai, *PLOS ONE* **10**, e0135286 (2015).
34. D. Majra, J. Benson, J. Pitts, J. Stebbings, *J. Infect.* **82**, 36–40 (2021).
35. Epoch Times, "[Exclusive] The secret of Wuhan Huanan Seafood Market testing" (Epoch Times, 2020); <https://www.epochtimes.com/gb/20/5/31/n12150755.htm>.
36. Wuhan Municipal Bureau of Landscape Architecture and Forestry, "Administrative penalties in 2019" (Wuhan Municipal Bureau of Landscape Architecture and Forestry, 2019); https://web.archive.org/web/2021117124950/http://ylj.wuhan.gov.cn/zwgl/zwxgkzl_12298/cfqz/xzcl/202011/12020110_1499879.shtml.
37. Y.-Z. Zhang, E. C. Holmes, *Cell* **181**, 223–227 (2020).
38. J. E. Pekar *et al.*, *Science* **377**, 960–966 (2022).
39. N. Chen *et al.*, *Lancet* **395**, 507–513 (2020).
40. Z. Li *et al.*, *Lancet Reg Health West Pac* **8**, 100094 (2021).
41. Z. He *et al.*, *Lancet* **397**, 1075–1084 (2021).
42. E. C. Holmes *et al.*, *Cell* **184**, 4848–4856 (2021).
43. J. Pekar, M. Worobey, N. Moshiri, K. Scheffler, J. O. Wertheim, *Science* **372**, 412–417 (2021).
44. L. Chang *et al.*, *Nat. Commun.* **12**, 1383 (2021).
45. L. Chang *et al.*, *Protein Cell* **wpac013** (2019).
46. L. Lu *et al.*, *Nat. Commun.* **12**, 6802 (2021).
47. H.-L. Yen *et al.*, *Lancet* **399**, 1070–1078 (2022).
48. M. Standaert, E. Dou, "In search for coronavirus origins, Hubei caves and wildlife farms draw new scrutiny," *The Washington Post*, 11 October 2021; https://www.washingtonpost.com/world/asia_pacific/china-covid-bats-caves-hubei/2021/10/10/082eb8b6-1c32-11ec-bea8-308ea134594f_story.html.
49. X.-D. Lin *et al.*, *Virology* **507**, 1–10 (2017).
50. Q. Li *et al.*, *N. Engl. J. Med.* **370**, 520–532 (2014).
51. B. Lin, M. L. Dietrich, R. A. Senior, D. S. Wilcove, *Lancet Planet. Health* **5**, e386–e394 (2021).
52. T. M. Davies, J. C. Marshall, M. L. Hazelton, *Stat. Med.* **37**, 1191–1221 (2018).
53. Data and code for: M. Worobey *et al.*, The Huanan Seafood Wholesale Market in Wuhan was the early epicenter of the COVID-19, Zenodo (2022); <http://doi.org/10.5281/zenodo.6786454>.

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SUPPLEMENTARY MATERIALS

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The Huanan Seafood Wholesale Market in Wuhan was the early epicenter of the COVID-19 pandemic

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Pandemic epicenter

As 2019 turned into 2020, a coronavirus spilled over from wild animals into people, sparking what has become one of the best documented pandemics to afflict humans. However, the origins of the pandemic in December 2019 are controversial. Worobey *et al.* amassed the variety of evidence from the City of Wuhan, China, where the first human infections were reported. These reports confirm that most of the earliest human cases centered around the Huanan Seafood Wholesale Market. Within the market, the data statistically located the earliest human cases to one section where vendors of live wild animals congregated and where virus-positive environmental samples concentrated. In a related report, Pekar *et al.* found that genomic diversity before February 2020 comprised two distinct viral lineages, A and B, which were the result of at least two separate cross-species transmission events into humans (see the Perspective by Jiang and Wang). The precise events surrounding virus spillover will always be clouded, but all of the circumstantial evidence so far points to more than one zoonotic event occurring in Huanan market in Wuhan, China, likely during November–December 2019. —CA

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