ABSTRACT: As the most active and top producing area of coalbed methane (CBM) in China, the southern Qinshui Basin (SQB) is dominated by anthracite. Due to the low permeability of coals, plenty of non-gas-producing and low production CBM wells exist in the SQB. The permeability enhancement through some technological means is the key to increasing the CBM production of this area. In this paper, some typical anthracites were selected from the Daning block of the SQB to assess the effect of acidification treatments on permeability enhancement. The maceral composition determination shows that approximately 15% of minerals exist in the collected coal samples, and the X-ray diffractometer (XRD) results reveal that the minerals consist primarily of clay minerals, along with a little amount of quartz, calcite, and dolomite. Two types of acidizing fluids were used to conduct acidification treatments on the anthracites for different lengths of time. The N₂ permeability of the anthracites before and after acidification was measured and compared. The results show that the original samples exhibit low permeability. As the acidification time increases, the permeability of all of the samples shows an increasing trend, and the acid sensitivity index $I_a$ increases rapidly first and then levels off, and finally approaches 1. After 48 h of acidification, the samples show an increase ranging from 8.75 to 22.67 times (avg. 14.3 times) the original permeability. The permeability enhancement of the SQB anthracites is mainly attributed to the dissolution of acid-soluble minerals in the cleat system of coal. The minerals in the cleats are completely or partially dissolved by the acids, generating some soluble and insoluble substances; when the fluid flows through, the cleat space is reallocated. Overall, the cleat demineralization by acids frees up a lot of cleat spaces, leading to an increase in cleat connectivity. As a result, the fluid movement becomes smooth and the permeability of coal improves.

INTRODUCTION

The commercial extraction of methane from coal beds is now well established in a number of countries throughout the world. The global coalbed methane (CBM) resources estimated by the International Energy Agency (IEA) amount to 263.8 trillion m³. The United States, Canada, Australia, China, and European countries such as Germany and the United Kingdom have all achieved the commercial development of CBM. Due to mine-safety considerations, as well as energy demand and environmental benefits, the CBM development has been attached great importance by the Chinese government in the past decade. According to the fourth resource evaluation results conducted by the Ministry of Land and Resources of China, the CBM resources of China are 30.05 trillion m³, and the recoverable resources are 12.5 trillion m³, ranking third in the world. Since 2008, the CBM production from surface wells of China has risen from 0.5 to 5.46 billion m³ in 2019.

The Qinshui Basin possesses approximately 3.28 trillion m³ CBM resources. As the largest and the most mature CBM development basin in China at present, the Qinshui Basin accounts for 80% of CBM production of China. The most successful CBM commercial development regions in the Qinshui Basin are in the southern portion, i.e., the southern Qinshui Basin (SQB). The maximum vitrinite reflectance ($R_{o,max}$) of coals in the SQB is between 2.0 and 4.0%, and mostly exceeds 3.0%, reaching the anthracite stage. Though more than 8000 surface CBM-producing wells have been drilled in the SQB, the total production is still unsatisfactory because the number of non-gas-producing and low production wells accounts for up to 50—75% of the total. Previous studies have shown that the key determinant for the gas production rate from a gas-bearing coal deposit is coal permeability. Therefore, the overall low permeability of...
high-rank coal is considered to be the vital reason for the low CBM production in the SQB.15−17

Due to low permeability, more than 96% CBM wells in China rely on fracturing stimulation technology to improve the permeability of coal seam.18 The routine fracturing technology applied to coal seam is hydraulic fracturing, which uses surface high-pressure pumps to force fracturing fluid containing sand particles into the coal seam at high pressure to cause the coal seam to crack and produce conductive fractures.19 The sand particles acting as a "proppant" are left in the coal seam as the fracturing fluid flows back to keep the newly formed fractures open and conductive. In general, about 98.5% of the hydraulic fracturing fluid is water and sand, and additives make up the rest 1.5%. The additives commonly consist of a variety of chemicals mainly aimed at structuring the fluid and assisting the delivery of the sand.20

The natural cleat system in the coal, including face cleat and butt cleat, plays a major role in determining permeability.1,14,21,22 Previous studies have discovered that the cleat system in the coal is usually mineralized to varying degrees by various minerals, which occlude cleat porosity and reduce the permeability of the coal. Highly mineralized cleats can lead to low permeability of coals.21,23−25 Hence, cleat demineralization by acids provides a possibility for improving coal permeability through increasing cleat connectivity and/or creating additional flow channels.19 Utilization of acidification treatments to enhance permeability has been applied to conventional sandstone26−28 and carbonate reservoirs.29−33 In recent years, this technology has been attempted to be applied to enhancing coal permeability.19,22,34−36

Many previous studies have verified that the natural fractures in the coals of the SQB are filled with various minerals, such as carbonate (e.g., calcite, dolomite), oxide (e.g., hematite, quartz), sulfide (e.g., pyrite), and silicate minerals (e.g., kaolinite, illite).17,37,38 The presence of these highly mineralized fractures significantly reduces the permeability of coals of the SQB. In this paper, some typical anthracites of the Daning block in the SQB were selected to assess the effect of acidification treatments on permeability enhancement. In addition, the mechanism of permeability enhancement of the SQB anthracite by acidizing was tentatively explored. The results can help improve the formulation of fracturing fluid applied to high-rank coals of the SQB.

■ GEOLOGICAL SETTINGS

The Daning block, covering an area of approximately 38.82 km², is located in the southern Qinshui Basin (Figure 1b), which is a tectonic basin formed on the basement of the Late Paleozoic era.17 The Daning block is situated at the junction of the southern end of Taihang Mountain and the northeastern edge of Zhongtiao Mountain.9 The stratum of the Daning block slopes from south to north. The dip of the stratum is generally within 10° and can be over 20° locally due to structural influence. The eastern part of the block is dominated by folds with axial striking NE−SW or near N−S; the western part develops faults striking NE−SW, NNE−SSW, or W−E (Figure 1c). The Sitou fault system constitutes the western boundary of the block and affects the gas content of the coal seam in the block (Figure 1c).

The strata in the study area include Ordovician Fengfeng formation (O2f), Carboniferous Benxi formation (C2b),
The thickness of no. 3 coal seam in this block varies from 2.21 to 6.97 m, averaging 4.45 m. The burial depth is approximately between 100 and 600 m. The maximum vitrinite reflectance \( R_{\text{omax}} \) ranges from 2.95 to 3.36\% (avg. 3.17\%), indicating that the coal in this region is typical anthracite. The well testing permeability of no. 3 coal seam is from 0.10 to 6.49 mD with an average value of 0.96 mD. The reservoir pressure obtained from well testing varies from 0.67 to 3.19 (avg. 1.62) MPa, and the reservoir pressure gradient ranges from 0.15 to 0.82 (avg. 0.43) MPa/100 m. The gas content (dry ash-free basis, \( G_{\text{daf}} \)) of no. 3 coal seam ranges from 5.56 to 17.57 m\(^3\)/t with an average of 12.83 m\(^3\)/t. The gas content inside the block is better than that near the boundary and is commonly low near the faults (Figure 1c).

![Figure 2](https://example.com/figure2.png)

**Figure 2.** Stratigraphic column of coal-bearing strata in the Daning block and the sampled section.

The thickness of no. 3 coal seam in this block varies from 2.21 to 6.97 m, averaging 4.45 m. The burial depth is approximately between 100 and 600 m. The maximum vitrinite reflectance \( R_{\text{omax}} \) ranges from 2.95 to 3.36\% (avg. 3.17\%), indicating that the coal in this region is typical anthracite. The well testing permeability of no. 3 coal seam is from 0.10 to 6.49 mD with an average value of 0.96 mD. The reservoir pressure obtained from well testing varies from 0.67 to 3.19 (avg. 1.62) MPa, and the reservoir pressure gradient ranges from 0.15 to 0.82 (avg. 0.43) MPa/100 m. The gas content (dry ash-free basis, \( G_{\text{daf}} \)) of no. 3 coal seam ranges from 5.56 to 17.57 m\(^3\)/t with an average of 12.83 m\(^3\)/t. The gas content inside the block is better than that near the boundary and is commonly low near the faults (Figure 1c).

**SAMPLES AND METHODS**

**Sample Collection and Preparation.** In this study, a total of three fresh coal samples (DN-1, DN-2, and DN-3) were collected from the underground coal mine in the Daning block. All of the three samples were collected from the no. 3 coal seam in Shanxi Formation (Figure 2). The collected coal samples were big blocks with approximate dimensions of 250 mm × 250 mm × 200 mm. To prevent oxidation and preserve the natural moisture content during transport, the coal samples were wrapped with a white preservative film and then placed in black plastic bags immediately after being removed from the working faces. Samples were then sent to the laboratory for sample preparation.

For each big block of sampled coal, a horizontal core was carefully drilled with an approximate diameter of 2.5 cm and length of 4–5 cm, parallel to the bedding plane of the coal sample. Some small block samples with approximate dimensions of 2 cm × 2 cm × 2 cm were taken from alongside the cores, and these were crushed and sieved to different sizes for use in other experiments: <1 mm for maceral group composition quantifications and \( R_{\text{omax}} \) measurements, 200 mesh for powder X-ray diffraction (XRD) and proximate analysis, and 10–20 mesh for static leaching experiments of coal fines.

**Proximate and Petrology Analysis.** Proximate analysis measurements of the coal samples including ash yield, moisture, and volatile matter were performed following the Chinese standard GB/T 212-2008. \( R_{\text{omax}} \) measurements and maceral group composition determination were performed using a Leitz MPV-3 photometer microscope, according to the Chinese standards of GB/T 6948-2008 and GB/T 8899-2013, respectively.

Mineral compositions of the coal samples were determined using a D8 DISCOVER X-ray diffractometer made by Bruker, Germany. Prior to XRD analysis, low-temperature ashing of the powdered coal samples was carried out using a YAMATO PR301 plasma asher. The XRD analysis procedure and semiquantitative analysis are described in detail in Li et al.39

**Static Leaching Experiments of Coal Fines.** In the oil field, the mixed acid of hydrofluoric acid (HF) and hydrochloric acid (HCl) is often referred to as “mud acid”, which is used to remove mud blockage and increase the permeability of mud and sandstone formations, thus facilitating water injection or oil production. HCl in the mud acid can dissolve the carbonate minerals and iron and aluminum compounds in the formations, and HF can dissolve the clay and silicate minerals. Considering that the reaction speed of HCl and HF is too fast, and CH\(_3\)COOH can slow down the dissolution speed, so a certain amount of CH\(_3\)COOH was added to the acid solution formula in this study. KCl acting as a stabilizer is a common additive in the fracturing fluid.

The static leaching experiments of coal fines were used to investigate the coal dissolution effect in different types of mixed acids. An appropriate amount of fresh block coal from the Daning block was chosen, crushed, and sieved into particles with sizes between 10–20 mesh. Then, the crushed coal particles were dried in a drying oven at 60 °C for 12 h. The dried coal particles were divided into 36 subsamples, and each subsample weighs 5 g, accurate to 0.001 g. Next, each subsample was soaked in 50 mL of different mixed acids (Table 1) for different lengths of time (Table 2) at ambient pressure (1 atm) and constant temperature of 30 °C, which
was provided by a thermostatic water bath. Then, the crushed coal particles and the leachate were separated using quantitative filter paper with a pore size of 30–50 μm. The coal fines along with the filter paper were dried at 80 °C until the mass is constant and weighed to determine the mass variation before and after acidizing. The dissolution rate ($R_d$) was calculated according to eq 1, as follows

$$R_d = \frac{w_1 - (w_f - w_0)}{w_1} \times 100\%$$

where $R_d$ is the dissolution rate of the coal, %; $w_1$ is the weight of the coal before acidizing, g; $w_f$ is the weight of the coal after acidizing, g; and $w_0$ is the weight of the filter paper, g.

The calculated dissolution rate of Daning coal in different types of mixed acids is presented in Table 2 and Figure 3. After acidizing for 48 h, the dissolution rate of Daning coal in mixed acids A, B, and C is the highest with values of 7.60, 7.80, and 7.90%, respectively. From the dissolution rate curves in Figure 3.

**Table 2. Dissolution Rate of Daning Coal Fines in Different Mixed Acids**

| time (h) | A (%) | B (%) | C (%) | D (%) | E (%) | F (%) |
|---------|-------|-------|-------|-------|-------|-------|
| 1       | 1.33  | 1.33  | 3.30  | 1.80  | 1.41  | 1.20  |
| 6       | 2.66  | 2.70  | 4.00  | 2.50  | 2.33  | 2.10  |
| 12      | 3.50  | 3.80  | 4.60  | 3.60  | 3.40  | 3.10  |
| 24      | 4.90  | 5.40  | 6.60  | 4.00  | 4.60  | 4.20  |
| 48      | 7.60  | 7.80  | 7.90  | 3.60  | 7.10  | 4.30  |

R is the N2 permeability of the coal sample, mD (10$^{-13}$ μm$^2$); $p_0$ and $p_1$ are gas pressures at the outlet and the inlet, MPa, respectively; $A$ and $L$ are the cross-sectional area, cm$^2$, and the length, cm, of the coal sample, respectively; $\mu$ is the viscosity of N2 under measured temperature, mPa-s; and $q$ is the flux of N2, cm$^3$/s.

**RESULTS AND DISCUSSIONS**

**Coal Properties and Mineral Compositions.** The measurement results of proximate analysis, maceral group compositions, and $R_{o,max}$ are presented in Table 3. Proximate analysis shows that moisture contents of the coal samples are between 0.92 and 2.11%, ash yields 10.11–15.16%, volatile matter yields 9.42–16.20%, and fixed carbon contents 83.80–90.58%, respectively. The maceral compositions data show that vitrinite contents account for 61.2–68.1%, inertinite contents between 16.8 and 17.6%, and mineral contents 14.9–16.2%.
Figure 4. Schematic diagram of the N$_2$ permeability experiment of the coal core.

Table 3. Coal Composition, $R_{o,max}$, and Proximate Analysis of the Coal Samples from the Daning Block

| coal samples | maceral and mineral (vol %) | $R_{o,max}$ (%) | proximate analysis (wt %) |
|--------------|----------------------------|------------------|--------------------------|
|              | $V$ | $I$ | $L$ | $M$ | $R_{o,max}$ | $M_{ad}$ | $A_d$ | $V_{daf}$ | $FC_{daf}$ |
| Dn-1         | 61.2 | 17.6 | 0   | 16.2 | 3.05   | 0.92    | 10.11 | 16.20 | 83.80 |
| Dn-2         | 67.5 | 16.8 | 0   | 15.7 | 2.95   | 2.11    | 15.16 | 13.09 | 86.91 |
| Dn-3         | 68.1 | 16.8 | 0   | 14.9 | 3.00   | 1.65    | 12.92 | 9.42  | 90.58 |

$^a$R$_{o,max}$ = the max vitrinite reflectance; $V$ = vitrinite; $I$ = inertinite; $L$ = liptinite; $M$ = mineral; $M_{ad}$ = moisture (air-dried basis); $A_d$ = ash (dry basis); $V_{daf}$ = volatile matter (dry, ash-free basis); and $FC_{daf}$ = Fixed carbon (dry, ash-free basis).

Figure 5. XRD spectra of coal samples from the Daning block. (a) Dn-1, (b) Dn-2, and (c) Dn-3.
respectively. \( R_{\text{comax}} \) of the coal samples ranges from 2.95 to 3.05\%, indicating that the coal samples are anthracites.

The XRD spectra of the coal samples are shown in Figure 5, and the interpreted mineral proportions are presented in Table 4. The results show that the mineral matters in the coals of the Daning block consist primarily of clay minerals, along with a little amount of quartz, calcite, and dolomite. For coal samples of Dn-1 and Dn-3, the clay minerals are composed of interstratified clay minerals, along with varying amounts of kaolinite, chlorite, and illite. However, the clay minerals in the coal sample of Dn-2 mainly consist of vast illite and relatively lower interstratified minerals.

### Changes of Permeability after Acidizing

The \( N_2 \) permeability of coal samples before and after acidifying is provided in Table 5.

#### Table 4. Mineralogical Proportions of the Coal Samples Determined by XRD from the Daning block (Unit in %)

| coal samples | quartz | calcite | dolomite | clay minerals | quartz | calcite | dolomite | kaolinite | chlorite | illite | interstratified minerals |
|--------------|--------|---------|----------|--------------|--------|---------|----------|-----------|----------|-------|-------------------------|
| Dn-1         | 4      | 2       | 1        |              | 22     | 9       | 2        | 60        |          |      |                         |
| Dn-2         | 1      | 3       | 1        |              | 1      | 0       | 65       | 29        |          |      |                         |
| Dn-3         | 2      | 1       | 1        |              | 5      | 4       | 29       | 58        |          |      |                         |

#### Table 5. Measurement Results of \( N_2 \) Permeability of the Coal Samples before and after Acidification\(^a\)

| coal samples | \( k_0 \) | acid type | \( k_0 \) | \( R_e \) | \( I_a \) | \( k_0 \) | \( R_e \) | \( I_a \) | \( k_0 \) | \( R_e \) | \( I_a \) |
|--------------|---------|-----------|---------|---------|---------|---------|---------|---------|---------|---------|---------|
| Dn-1         | 0.45    | A         | 1.45    | 3.22    | 0.69    | 3.02    | 6.71    | 0.85    | 10.2    | 22.67   | 0.96    |
| Dn-2         | 1.21    | A         | 3.26    | 2.69    | 0.63    | 8.32    | 6.88    | 0.85    | 13.97   | 11.55   | 0.91    |
| Dn-3         | 0.12    | B         | 0.16    | 1.33    | 0.25    | 0.21    | 1.75    | 0.43    | 1.05    | 8.75    | 0.89    |

\(^a\) \( k_0 \) \( N_2 \) permeability of coal sample before acidification, mD; \( k_0 \) \( N_2 \) permeability of coal sample after different acidification length of time, mD; \( R_e = k_a/k_0 \); and \( I_a \), acid sensitivity index.

#### Figure 6. Plots of the permeability (a) and acid sensitivity index (b) of coal samples versus acidification length of time.

#### Figure 7. Ratio change of \( k_a/k_0 \) of coal samples with acidification length of time.
presented in Table 5. The permeability of coal samples \( k_0 \) before acidification shows a low level with values ranging from 0.12 to 1.21 mD, averaging only 0.59 mD. \( k_0 \) of the three coal samples takes on \( D_{n-2} > D_{n-1} > D_{n-3} \). After acidizing the coal samples for different lengths of time, the permeability of coal samples \( k_a \) was increased to varying degrees. It can be seen that the permeability of all of the samples shows an increasing trend as the acidification time increases. That is, the longer the acidification time, the greater the increase in permeability. Within the same acidification time, the permeability value is always maintained at \( D_{n-2} > D_{n-1} > D_{n-3} \) (Table 5 and Figure 6a). After 48 h of acidification, sample \( D_{n-1} \) presents the largest increase in permeability, which increases from 0.45 to 10.2 mD, an increase of 21.67 times the original permeability; samples \( D_{n-2} \) and \( D_{n-3} \) have an increase of nearly 10 times (Figure 7). Obviously, acidification can significantly improve the permeability of anthracites of the SQB. In terms of acid type, type A acid is better than type B acid in improving the permeability of the coal samples (Table 5).

To characterize the sensitivity of coal reservoir permeability to acid treatments, a parameter named the acid sensitivity index \( I_a \) defined in this study, and the calculation formula is as follows

\[
I_a = \frac{k_a - k_0}{k_0}
\]

where \( k_a \) is the \( N_2 \) permeability of the coal sample after acidification, mD; \( k_0 \) is the \( N_2 \) permeability of the coal sample before acidification, mD; and \( I_a \) is the acid sensitivity index of the coal sample.

It can be seen that the acid sensitivity index \( I_a \) increases with the increase in acidification time (Figure 6b). The \( I_a \) of samples \( D_{n-1} \) and \( D_{n-2} \) is always maintained at the same level, and \( I_a \) of sample \( D_{n-3} \) is significantly lower than the other two samples at most of the acidification times. It reflects that the permeability of sample \( D_{n-3} \) is relatively insensitive to acid treatment. Overall, with the increase in acidification time, \( I_a \) increases rapidly first and then levels off, and finally approaches 1.

**Mechanism of Permeability Enhancement by Acidizing.** A coal reservoir is a dual-porosity system containing porous matrix blocks and a naturally fractured network known as cleats.\(^{40-43}\) The complex natural cleats are widespread in coal reservoirs and play a crucial role in the permeability of coal reservoirs. However, lots of cleats in coal reservoirs are filled with various minerals such as carbonate (e.g., calcite, dolomite), oxide (e.g., hematite, quartz), sulfide (e.g., pyrite), and silicate minerals (e.g., kaolinite, illite).\(^{24,25}\) The presence of these highly mineralized cleats may occlude fracture porosity and then reduce the permeability of coal reservoirs.\(^{19}\)

As shown in Tables 3 and 4, the anthracites in this study contain approximately 15% minerals, which are primarily composed of clay minerals, along with a little amount of quartz, calcite, and dolomite. The previous studies in the SQB have verified that these minerals widely exist in the cleats and pores of anthracite (Figure 8).

![Figures](https://doi.org/10.1021/acsomega.1c04539)
During the acidification process, a large amount of acids (HCl, HF) enters the pores and cleats of the coal reservoirs and then reacts with the minerals. The possible chemical reaction equations involved are presented in eqs 3–7. Carbonate minerals are converted into water-soluble chlorides under the action of HCl (eqs 3 and 4). Quartz is converted into fluorosilicic acid under the slow dissolution of HF (eq 5). Kaolinite reacts with HF to generate fluorosilicic acid and insoluble aluminum fluoride (eq 6). Smectite reacts with HF to generate fluorosilicic acid and fluoroaluminate (eq 7).

Acidification of carbonate minerals

calcite: CaCO₃ + 2HCl → CaCl₂ + CO₂ + 2H₂O

(3)
dolomite: CaMg(CO₃)₂ + 4HCl
→ CaCl₂ + MgCl₂ + 2CO₂ + 2H₂O

(4)

Acidification of quartz
quartz: SiO₂ + 6HF → H₄SiF₆ + 2H₂O

(5)

Acidification of clay minerals
kaolinite: Al₂Si₂O₅(OH)₄ + 18HF
→ 2AlF₃ + 2H₂SiF₆ + 9H₂O

(6)
smectite: Al₂O₃·4SiO₂·2H₂O + 36HF
→ 2H₄AlF₆ + 4H₂SiF₆ + 12H₂O

(7)

The graphical representation showing the dissolution of acid-soluble minerals in cleats and pores is presented in Figure 9. As the channels of fluid flow, the cleats (face cleat and butt cleat) distributed between the coal matrix blocks were completely or partially filled with large amounts of minerals before acidification of coal samples. These minerals, including clay minerals, quartz, calcite, and dolomite, block the cleat space and seriously affect the movement of fluids. The anthracites in the study area have poor in situ cleat development and low permeability. The presence of these highly mineralized cleats will undoubtedly lead to much lower permeability. With the action of acids, the minerals in the cleats are completely or partially dissolved. The carbonate minerals were completely dissolved and converted into water-soluble chlorides, which were then carried away by the fluid flow. The quartz and clay minerals are partially dissolved, and it causes them to break up into small particles, among which some remain on the wall of the cleats and some become mobile and are carried away by the fluid flow. During the dissolution of clay minerals by acids, some insoluble substances such as aluminum fluoride (AlF₃) were generated and accumulated on the surface of the lower wall of the cleat or carried away by the fluid flow. Overall, the cleat demineralization by acids frees up a lot of cleat spaces, leading to an increase in cleat connectivity. As a result, the fluid movement becomes smooth and the permeability of coal improves. During the acidification, some minerals in the pores of the matrix will also be dissolved by acids, which will improve the gas diffusion of coal.

CONCLUSIONS

Mineral occlusions in cleats considerably reduce the permeability of anthracites in the SQB, influencing CBM production. In this work, some anthracites from the SQB were selected to assess the effect of acidification treatments on permeability enhancement. For all of the samples, an obvious increasing trend of permeability was obtained as the acidification time increased. The samples show an increase ranging from 8.75 to 22.67 times (avg. 14.3 times) the original permeability after 48 h of acidification. Overall, the acid sensitivity index Iₐ increases rapidly first and then levels off, and finally approaches 1. The permeability enhancement of the SQB anthracites is mainly attributed to the dissolution of acid-soluble minerals in the cleat system of coal. The minerals in the cleats are completely or partially dissolved by the acids, generating some soluble and insoluble substances; when the fluid flows through, the cleat space is reallocated. Overall, the cleat demineralization by acids frees up a lot of cleat spaces, leading to an increase in cleat connectivity. As a result, the fluid movement becomes smooth and the permeability of coal improves. Acidification treatments have an obvious effect on increasing the permeability of anthracites in the SQB.
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Notes
The authors declare no competing financial interest.

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