Supplementary Figure 1. Stable carbon isotope data at 391 m water depth (in °/oo with respect to PDB standard). (a) δ^{13}C values of dissolved inorganic carbon (DIC). (b) δ^{13}C values of dissolved methane.

Supplementary Figure 2. Isotopic composition of pore fluids versus dissolved chloride concentration at 391 m water depth. Lines are calculated by mixing ambient seawater with freshwater from different sources including meltwater from northwest Svalbard (meteoric), water released form clay minerals during sediment diagenesis (clay), and hydrate water released upon gas hydrate dissociation (hydrate). (a) δ^{18}O values of pore water with respect to Vienna Standard Mean Ocean Water (VSMOW). (b) δ^{2}H values of pore water with respect to VSMOW.
Supplementary Figure 3. Molar Li/Cl and B/Cl ratios in pore fluids. (a) Li/Cl at 391 m water depth. (b) B/Cl at 391 m water depth. (c) Li/Cl at 404 m water depth. (d) B/Cl at 404 m water depth.
Supplementary Figure 4. Model results for hydrate melting at 391 m water depth. The first row (a, b, c) shows the initial values at 1980 and final results for 2016 of the simulation depicted in Fig. 4. The second row (d, e, f) shows the corresponding results for a different choice of the initial hydrate depth profile. The third row (g, h, i) depicts simulation results for elevated initial temperatures and hydrate saturations.
Supplementary Figure 5. Model results for Holocene hydrate melting at 391 m water depth. (a) Bottom water temperatures ($T_{BW}$) applied as model forcing. (b) Percent of pore space occupied by gas hydrate ($Sat_{GH}$). (c) Bulk sediment temperature ($T$). Dots indicate temperatures measured in drill holes at 391 m water depth (s. Fig. 2). (d) Dissolved chloride concentration in pore fluids ($Cl$). Dots indicate concentrations in cores retrieved at 391 m water depth (s. Fig. 2).
Supplementary Figure 6. Eustatic sealevel (solid line) and change in seabed elevation at the upper continental slope of northwestern Svalbard as calculated with the ice sheet model (broken line). Relative sea level change is calculated as difference between eustatic sealevel and seabed elevation.

Supplementary Figure 7. Model results for sediment temperature. (a) 391 m water depth. (b) 404 m water depth. The bottom water temperature was maintained at 2.5°C for 8 ka – 0.1 ka and was enhanced to 3°C over the last 100 years of the model period. Temperature gradients of 45°C km\(^{-1}\) (391 m) and 50°C km\(^{-1}\) (404 m) were applied at the base of the model column (100 mbsf). Dots indicate data in cores retrieved at 391 m and 404 m water depth. Final temperature profiles at 0 ka are also shown in Fig. 2.
**Supplementary Table 1**: Location of the coring and drilling sites

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<th>Water depth (m)</th>
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**Supplementary Discussion**

The freshwater component in Cl-depleted pore fluids could be meteoric water originating from Svalbard, water released from smectite and other water-bearing mineral phases during sediment diagenesis, a salt-depleted fluid created during hydrothermal phase separation, or water released during the dissociation of gas hydrates\(^2\). The stable isotopic composition of pore water (δ\(^{18}\)O, δ\(^3\)H) indicates that the freshwater component is neither of meteoric origin nor produced by the dewatering of smectite and other clay minerals (Supplementary Figure 2). The most plausible meteoric source would be glacial meltwater from northwestern Svalbard with a mean δ\(^{18}\)O\(_m\) value of -14.7 °/oo \(^3\) and a δ\(^3\)H\(_m\) value of about -100 °/oo as derived from the local meteoric water line\(^4\). Clay minerals formed by weathering processes on Svalbard in equilibrium with the depleted meltwater should contain chemically bound water enriched in δ\(^{18}\)O and further depleted in δ\(^3\)H \(^5\) with a composition of δ\(^{18}\)O\(_c\) = +14.2 °/oo and δ\(^3\)H\(_c\) = -137 °/oo \(^6\). Mixing between these freshwater endmembers (meteoric, clay-derived) and near surface porewater (Cl = 556 mM, δ\(^{18}\)O\(_w\) = 0.3 °/oo, δ\(^3\)H\(_w\) = 2.1 °/oo) yields isotopic compositions for the Cl-depleted fluids that strongly deviate from our data (Supplementary Figure 2). Dissolved Li and B are depleted in all porewater samples with respect to seawater (Supplementary Figure 3). Li and B are removed from pore fluids at low temperatures\(^7,\(^8\) and released into the pore water when sediments are heated to more than about 60°C\(^9\). Since the depleted dissolved B and Li values suggest a low-temperature origin of the freshened fluids, the freshwater does not originate from hydrothermal phase separation and mineral dewatering processes such as smectite-illite transformation that typically occurs at >60°C\(^10\).

Since all other freshwater sources can be excluded, we conclude that the chloride depletion is caused by in-situ gas hydrate dissociation. Gas hydrates are enriched in both \(^{18}\)O and \(^2\)H with respect to ambient porewater\(^11\). At our sites, the water bound in methane hydrate should have isotopic compositions of δ\(^{18}\)O\(_h\) = 2.6 – 3.5 °/oo and δ\(^3\)H\(_h\) = 16 – 24 °/oo \(^11\). Dissociation of gas hydrates with δ\(^{18}\)O\(_h\) = 2.6 °/oo and δ\(^3\)H\(_h\) = 16°/oo yields a mixing line that is broadly consistent with the data (Supplementary Figure 2). However, most samples tend to plot below the hydrate line even though we applied minimum estimates for hydrate isotope values. These second order deviations may be explained by the precipitation of \(^{18}\)O-enriched carbonates and the microbial oxidation of organic matter and methane that releases \(^2\)H-depleted water into the pore space\(^5\). It is also possible that water cages of the dissociating gas hydrates were not completely occupied by gas molecules and therefore even less enriched in \(^{18}\)O and \(^2\)H than assumed in our calculation\(^12\). Finally, the low isotopic fractionation observed in the Cl-depleted pore fluids may be explained by diffusion. The molecular
diffusion coefficient of water is higher than that of Cl and Na such that $\delta^{18}$O and $\delta^2$H enrichments may relax more rapidly to background values than chloride and salinity depletions.

**Supplementary References**


