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Article

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Hydrogen (H₂) is expected to play a crucial role in reducing greenhouse gas emissions. However, hydrogen losses to the atmosphere impact atmospheric chemistry, including positive feedback on methane (CH₄), the second most important greenhouse gas. Here we investigate through a minimalist model the response of atmospheric methane to fossil fuel displacement by hydrogen. We find that CH₄ concentration may increase or decrease depending on the amount of hydrogen lost to the atmosphere and the methane emissions associated with hydrogen production. Green H₂ can mitigate atmospheric methane if hydrogen losses throughout the value chain are below $9 \pm 3\%$. Blue H₂ can reduce methane emissions only if methane losses are below 1%. We address and discuss the main uncertainties in our results and the implications for the decarbonization of the energy sector.

Commitments to reach net-zero carbon emissions have drawn renewed attention to hydrogen (H₂) as a low-carbon energy carrier^{1,2}. Currently, H₂ is mostly used as an industrial feedstock, and its global production has a high carbon footprint because it relies almost entirely (\approx 95%) on fossil fuels¹. However, many technologies to produce H₂ with a lower carbon footprint are available¹. Among these, low-carbon H₂ can be produced from water electrolysis powered by renewable energy (green H₂) or from methane reforming coupled with carbon capture and storage (blue H₂). H₂ fuel may be especially important to decarbonize energy and transport sectors where direct electrification is complicated, like heavy industry, heavy-duty road transport, shipping, and aviation¹. H₂ is also being considered for storing renewable energy¹. As a result of this potential, countries accounting for more than a third of the world's population have developed national strategies for large-scale H₂ production^{1,2}.

Even if a more hydrogen-based economy would reduce CO_2 emissions and improve air quality³, it would also increase the H_2 emissions into the atmosphere. The H_2 molecule is very small and difficult to contain, so it is still largely unknown how much H_2 will leak in future value chains. H_2 emissions will also occur due to venting, purging, and incomplete combustion^{4–6}. This potential increase in H_2

emissions has received relatively little attention to date because H₂ is neither a pollutant nor a greenhouse gas (GHG). However, it has been long known⁷⁻¹⁰ that H₂ emissions may exert a significant indirect radiative forcing by perturbing the concentration of other GHG gases in the atmosphere. This indirect GHG effect of H₂ calls for a detailed scrutiny of the global H₂ budget and the environmental consequences of its perturbation^{11,12}.

 H_2 is the second most abundant reactive trace gas in the atmosphere, after methane, with an average concentration of around 530 ppb_v¹³. H₂ sources include both direct emissions (≈45% of total sources) and production in the troposphere from the oxidation of volatile organic compounds (≈25%) and methane (≈30%)^{11,14}. The main H₂ sinks are the uptake by soil bacteria (70–80% of total tropospheric removal) and the atmospheric reaction with the radical OH (20–30%), which is responsible for the indirect GHG effect of H₂. H₂'s reaction with the OH radical tends to increase tropospheric methane (CH₄) and ozone (O₃), which are two potent greenhouse gases. It also increases stratospheric water vapor, which is associated with stratospheric cooling and tropospheric warming^{8,15}. Recent global climate models have estimated that hydrogen has an indirect radiative forcing of around 1.3¹⁴–1.8¹⁶ 10⁻⁴ W m⁻² ppb_v⁻¹, and a global warming potential (GWP) that lies in the

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Fig. 1 | **Tangled hydrogen (H₂) and methane (CH₄) budgets.** Sketch of H₂ and CH₄ tropospheric budgets and their interconnections: (1) the competition for OH; (2) the production of H₂ from CH₄ oxidation; (3) the potential emissions [minimum-maximum] due to a more hydrogen-based energy system. Flux estimates (Tg/year) are from refs. 11,18. Arrows are scaled with mass flux intensity, CH₄ scale being 10

times narrower than H₂ scale. On a per-mole basis, H₂ consumes only around 3 times less OH than CH₄. ppq = part per quadrillon (10⁻¹⁵). ^{*a*} top-down estimate including also minor atmospheric sinks (<10%). ^{*b*} range obtained as a difference between total and fossil fuel emissions¹⁸.

range 11 ± 5 for a 100-year time horizon¹⁶. Hence, H₂ emissions are far from being climate neutral, and their largest impact is related to the perturbation of atmospheric $CH_4^{14,16}$, the second most important anthropogenic GHG.

The tropospheric budgets of H₂ and CH₄ are deeply interconnected (Fig. 1). First, the removal of both gases from the atmosphere is controlled by their reaction with OH, which is the dominant sink (\approx 90%) for atmospheric methane^{17,18}. An increase in the concentration of tropospheric H₂ may reduce the availability of OH, consequently weakening CH₄'s removal and increasing CH₄'s lifetime and abundance^{14,19}. Second, methane is a primary precursor of hydrogen. Namely, CH₄ oxidation results in the production of formaldehyde, whose photolysis produces H₂. Firn-air records suggest that the increase in H₂ over the 20th century can be largely explained by the increase in CH₄ concentration²⁰.

Additionally, H_2 and CH_4 are linked at the industrial level. Around 60% of global H_2 production is currently produced from steam methane reforming (gray H_2) and is responsible for 6% of global natural gas use¹. In the next decade, steam methane reforming coupled with carbon capture and storage will likely remain the dominant technology for large-scale H_2 production (blue H_2), since facilities for H_2 production from renewable sources (green H_2) will require time to become operational and economically favorable².

Since CH_4 is the second-largest contributor to atmospheric warming since the beginning of the industrial era and there are global efforts to mitigate its atmospheric levels²¹, it is crucial to quantify the response of atmospheric CH_4 to increasing H_2 production.

We analyze this problem through a simple atmospheric model that captures the interaction between H₂ and CH₄ ("Methods"). The investigation of the transient dynamics ("Methods") shows that any H₂ emissions pulse to the atmosphere leads to a small transient growth of atmospheric CH₄ whose effects last for several decades. In the next sections, we focus on how the equilibrium concentrations of tropospheric H₂ and CH₄ would respond to scenarios of continuous emissions from an energy system where part of the fossil fuel energy share is replaced by green or blue H₂. The analysis emphasizes how atmospheric CH₄ could either decrease or increase, mainly depending on the H₂ production pathway and the amount of H₂ lost to the atmosphere. The latter is defined through the hydrogen emission intensity (HEI), namely the percentage of H₂ produced that is lost to the atmosphere. Specifically, we find a critical HEI above which the CH_4 atmospheric burden rises despite the lower fossil fuel use. We assess the critical factors and the main uncertainties in the quantification of this critical HEI. We finally discuss how our results can help better inform policymakers regarding the trade-off associated with different scenarios of hydrogen production and use.

Results

Emission scenarios

Here we investigate how the tropospheric burdens of methane and hydrogen would be affected by the transition to a more hydrogenbased energy system, wherein hydrogen replaces part of the current fossil fuel energy (\approx 490 ExJ in 2019²²). To achieve this goal, we estimate the CH₄ and H₂ source changes, ΔS_{CH_4} and ΔS_{H_2} , where Δ indicates the difference to the current tropospheric conditions ("Methods"). This fossil fuel displacement reduces both CH₄ and H₂ sources (Fig. 1). The rise in H₂ production causes additional H₂ emissions due to intentional (e.g., venting) and unintended (e.g., fugitive) losses, and possibly CH₄ emissions associated with blue H₂ production.

The change in H₂ emissions can be estimated from the amount of hydrogen produced to substitute fossil fuels and the HEI, namely the percentage of H₂ produced that is lost to the atmosphere. Losses can occur due to venting, purging, incomplete combustion and leaks across the hydrogen value chain. The HEI of the future global H₂ value chain is very uncertain. Literature values range from 1 to 12%^{4,9,23}, but the upper bound is unlikely to occur at large scales because it would be both unsafe and too expensive. Recent empirical estimates for specific H₂ infrastructures suggest HEI's ranging from 0.1 to 6.9%, critically depending on the pathway of hydrogen production and transport⁶. To account for these uncertainties and to explore a broad spectrum of possible scenarios, here we vary HEI from 0 to 10% of the total hydrogen produced (Fig. 2a). The lower and upper bounds of this range represent a perfectly sealed and a highly leaking global H₂ value chain, respectively. With a perfectly sealed hydrogen value chain, H₂ emissions would only decrease due to the lower fossil fuel use. On the contrary, a highly leaking H_2 value chain, coupled with an envisioned penetration of H_2 in the energy market, could increase hydrogen emissions up to several times the total current sources, which are around 80 Tg H_2 yr⁻¹.

The variation in CH_4 emissions depends not only on the percentage of fossil fuel energy that is displaced by hydrogen, but also on the



Fig. 2 | **Hydrogen replacement of fossil fuels. a** Changes in H₂ sources (ΔS_{H_2}) as a function of fossil fuel replacement for different hydrogen emission intensity (HEI). **b** Changes in CH₄ sources (ΔS_{CH_4}) as a function of fossil fuel replacement for different H₂ production pathways. Methane leak rates associated with blue H₂ production are 0.2, 1, and 2%. Bands for ΔS_{CH_4} account for different amounts of blue H₂ produced and lost. **c** Response of the tropospheric concentrations of H₂

and CH₄ for the emission scenarios of the previous panels. Symbols mark the different percentages of fossil fuel displacement. Only symbols for 100% fossil fuel replacement are reported for blue H₂ with 1% CH₄ leakage. Also reported is the difference in CO₂ concentration (Δ [CO₂e]) that would produce equivalent radiative forcing to the change in equilibrium CH₄ (upper axis).

hydrogen production pathway. For green H₂, i.e., hydrogen obtained from renewable sources, we scale CH₄ emissions based on the reduced consumption of fossil fuels resulting from hydrogen usage (Fig. 2b). Estimates of current methane emissions associated with fossil fuel extraction and distribution are in the range 80–160 Tg CH₄ yr^{-118,24,25} and relatively equally distributed among coal, oil, and gas sectors²⁶. Here we use the top-down estimate of 111 Tg/year¹⁸.

For blue H₂, which is derived from steam methane reforming (SMR), the variation in CH4 sources not only accounts for the reduced consumption of fossil fuels but also for the methane emissions (venting, incomplete combustion, fugitive) associated with blue hydrogen production. These emissions depend on the amount of CH₄ needed to produce H₂, i.e., feedstock and energy requirements of the SMR process ("Methods"), and the CH₄ leak rate. The precise average leak rate of the global natural gas supply chain remains uncertain. One of the reasons is that national inventories generally underestimate real emissions²⁷⁻³⁰. More detailed studies relying on field measurements in the United States and Canada estimate average leak rates around 2%²⁸⁻³⁰, with large spatial heterogeneity between different operators³¹. Although national inventories suggest that some countries, like Venezuela and Turkmenistan, have higher leak rates²⁶, here we adopt 2% as the maximum global CH₄ leak rate for our scenarios, because methane-mitigation efforts are likely to decrease future global leak rates²¹ and, more importantly, because not all hydrogen produced will be blue H₂. In this regard, the scenario of blue H₂ with a 2% CH₄ leak rate can also be interpreted as a combination of equal production of green H₂ and blue H₂ with 4% CH₄ leak rate. We use 0.2% as a lower bound for the CH₄ leak rate, since this has been declared as the target of several energy companies for 2025^{32} . 1% represents an intermediate scenario of blue H₂ production.

Figure 2b shows the resulting CH₄ emissions associated with green and blue H₂ production with methane leak rates of 0.2, 1, and 2%. The different leak rates have a great impact on the methane emissions. Compared to the fossil fuel energy system, CH₄ emissions are reduced in the blue H₂ scenario with 0.2% methane losses, but largely increased in the blue H₂ scenario with 2% methane losses. The fossil fuel displacement by blue H₂ with 1% methane losses shows basically no net effect on the CH₄ emissions.

As a specific case, we also investigate the H_2 and CH_4 emission changes associated with estimates of future hydrogen production in a set of net-zero scenarios. H_2 production is expected to increase from current 90 Tg/year to 530–660 Tg/year in 2050^{2,33,34}. We thus consider a 500 Tg/year rise in the global H_2 production, which is energetically equivalent to about 15% of current fossil fuel energy. Figure 3a shows how, depending on the H_2 production pathway and the different hydrogen and methane leak rates, the emission changes of these two gases can vary substantially.

Tropospheric response

For the previous emission scenarios, we evaluate the changes in the equilibrium concentrations of tropospheric hydrogen and methane, namely Δ [H₂] and Δ [CH₄]. The timescales to equilibrium are dictated



Fig. 3 | Methane response to increasing H₂ production. a Changes in H₂ and CH₄ sources (ΔS) due to green and blue H₂ production (\approx 500 Tg yr⁻¹). HEI is the H₂ emission intensity. Gray lines mark the case for HEI = 0%. Blue bars for ΔS_{CH_2} are



obtained with HEI = 10%. **b** Response of CH₄ atmospheric concentration. The right axis shows the Δ [CO₂e] that would produce equivalent radiative forcing to the change in equilibrium CH₄.

by the gas average lifetimes ("Methods"). The corresponding variations in steady state concentration of OH are reported in Supplementary Figs. 1 and 2.

The H₂ economy causes a rise in tropospheric H₂ as a result of the additional emissions (Fig. 2c). The intensity of this increase varies considerably as a function of the emissions of the hydrogen value chain. The concentration variation could go from less than 100 ppb_v to more than 2000 ppb in envisioned scenarios of the H₂ economy, namely a +300% from the current H₂ tropospheric level.

The response of atmospheric CH₄ results from the combination of the methane emission change and the methane sink weakening due to the higher hydrogen emissions. To discriminate between the two mechanisms, it is useful to focus on the scenarios of fossil fuel displacement by green H₂. In the case of a perfectly sealed green H₂ value chain (HEI = 0%), [CH₄] and [H₂] both decrease due to reduction in fossil fuel emissions. As H₂ emissions increase (HEI > 0), Δ [CH₄] increases too. Up to the point that when HEI overcomes a critical threshold, there is an increase in atmospheric methane, i.e., Δ [CH₄] > 0, even though methane emissions are lower. This critical HEI is in the range 8–10% for green H₂ as it has a weak nonlinear dependence on the percentage of fossil fuel energy that is replaced by H₂ (see also Supplementary Fig. 3).

The scenarios of blue H_2 with 0.2% CH_4 leak rates are not very different from the green H_2 scenarios, with the critical HEI being in the range 7–8%. Regarding the scenarios of blue H_2 with 1% CH_4 leak rates, since there is basically no change in the methane emissions (Fig. 2b), the methane response is only associated with the reduction in OH availability due to the higher H_2 concentration. The critical HEI is not defined for this blue H_2 as the methane burden increases in all cases. The worst scenarios of blue H_2 with 2% CH_4 leak rates show drastic differences in the tropospheric concentrations of the two gases, which increase considerably, with a weakly nonlinear effect due to the drop in atmospheric OH.

The atmospheric methane response to future H₂ production^{2,33,34} shows qualitatively similar results as a function of the H₂ production pathway and the percentage of H₂ lost to the atmosphere (Fig. 3b). Positive effects in terms of methane mitigation are observed only for green and blue H₂ with low methane losses, if the H₂ emission intensity is well below 10%. Otherwise, the tropospheric methane burden is enhanced.

We also evaluated the change in CO₂ concentration (Δ [CO₂e]) that would produce equivalent radiative forcing to the change in the equilibrium concentration of CH₄ (Figs. 2c and 3b). We used the radiative efficiency of CH₄ that includes indirect effects on O₃ and stratospheric H₂O³⁵. Under the worst scenario of blue H₂ production with 2% CH₄ losses and 10% H₂ losses, the rise in equilibrium CH₄ due to future H₂ production would be like adding 9 ppm of CO₂ to the atmosphere (Fig. 3b). For the same blue H₂, the rise in CH₄ following the entire displacement of fossil fuels would be like adding around 70 ppm of CO₂ (Fig. 2c). This is equivalent to around 50% of the CO₂ increase from preindustrial times (278 ppm) to current days (417 ppm). Since the goal of keeping the global average temperature rise below 1.5 °C requires a mid-century maximum of CO₂ close to 450 ppm, these results support previous concerns about the sustainability of blue H₂³⁶ unless fugitive emissions can be kept sufficiently low.

Critical HEI for methane mitigation

The quantification of the critical hydrogen emission intensity (HEl_{cr}) for methane mitigation is key to assess whether displacing fossil fuels with hydrogen would mitigate or enhance the tropospheric burden of CH₄. Here we investigate how the HEl_{cr} is affected by the hydrogen production pathway and by two of the most uncertain terms in the CH₄-H₂-OH balance: (i) the partitioning of the OH sink among the tropospheric gases; (ii) the rate of H₂ uptake by soil bacteria. The derivation of an analytical solution for the HEl_{cr} is reported in the "Methods".

The very short lifetime of OH makes the quantification of its atmospheric dynamics extremely challenging. Indirect methods are typically used to estimate OH concentrations, sources, and sink partitioning³⁷⁻³⁹. Using a range of OH partition estimates^{38,40}, we investigate the dependence of the HEl_{cr} to different values of OH excess (E_{OH}), E_{OH} being the excess of OH that is consumed by other tropospheric gases besides hydrogen, methane, and carbon monoxide. Figure 4 shows the quasi-linear response of the HEl_{cr} to E_{OH} . We stress that a variation in E_{OH} is equivalent to a variation in the OH sources since we preserve the current average OH concentration, which is relatively well constrained by inverse modeling^{37,41}.

The HEI_{cr} is much lower for blue H₂ than for green H₂ because of the methane emissions associated with blue H₂ production. For the current tropospheric conditions, we find that HEI_{cr} is around 9% for green H₂, around 7% for blue H₂ with 0.2% methane leak rates, and 4.5% for blue H₂ with 0.5% methane leak rates. Blue H₂ with 1% methane leak rate has a HEI_{cr} that is close to zero, as displacement of fossil fuel with this hydrogen does not reduce methane emissions (Fig. 3b). For even higher methane leak rates, the methane burden would increase regardless of the H₂ emissions, so that the HEI_{cr} is negative.

The H₂ uptake by soil bacteria is another crucial process in the evaluation of HEI_{cr} and in the overall CH_4 – H_2 –OH dynamics, since it accounts for 70–80% of H₂ tropospheric removal¹¹. Despite recent research on uptake modeling^{42,43} and the microbial characterization of the H₂-oxidizing bacteria⁴⁴, the spatial heterogeneity of the uptake as driven by local hydro-climatic and biotic conditions hinders bottom-up estimates of the global average uptake rate. In atmospheric studies, the average uptake rate is usually adjusted in order to obtain a reasonable simulation of observed surface hydrogen concentrations^{14,45}.



Fig. 4 | **Critical hydrogen emission intensity (HEI) for methane mitigation.** Critical HEI as a function of OH excess (E_{OH}) and hydrogen production method (green and blue H₂ with 0.2, 0.5, 1% CH₄ leak rates, respectively). Dashed (dotted) lines are obtained for a 20% increase (decrease) in the H₂ uptake rate by soil bacteria (k_{cl}). Triangles mark the critical HEI for the best estimate of E_{OH} .

To account for these potential sources of uncertainties, we show how a $\pm 20\%$ variation in the uptake rate influences the critical HEI (bands in Fig. 4). A stronger biotic sink (dashed lines) reduces the consumption of OH by H₂ and, consequently, increases the HEI_{cr}. A weaker biotic sink (dotted line) has the opposite effect.

Regarding the impact of climate change on the H₂ soil sink, recent studies indicate that increasing temperatures are expected to slightly favor the uptake on a global scale¹⁴, while shifts in rainfall regimes will be the significant drivers of H₂ uptake changes at the local scale⁴³. From a biotic perspective, the adaptability of H₂-oxidizing bacteria to extreme environments⁴⁶ suggests that their presence will remain widespread in the future, but their spatial heterogeneity may change as a result of climate and anthropogenic pressures.

Another source of uncertainties in the evaluation of HEI_{cr} is related to the estimate of CH₄ emissions associated with fossil fuel use. Since there is a quasi linear relationship between these emissions and the HEI_{cr} (Eq. (16) in "Methods"), the same relative uncertainty of fossil fuel methane emissions (Fig. 1) applies to the HEI_{cr} .

Discussion

The success of the global net-zero transition hinges on hydrogen as a scalable low-carbon energy carrier that can replace fossil fuels in several hard-to-electrify energy and transport sectors. More than 20 governments and many companies have already announced strategies for hydrogen production, and the numbers are likely to increase as policy frameworks that facilitate hydrogen adoption are promoted^{1,2}. Considerable investments are still needed to achieve such a transition, as the current hydrogen momentum falls short compared to net-zero goals. The Hydrogen Council² estimates that there is a USD 540 billion gap between the investments of announced projects (USD 160 billion) on hydrogen production and the investments required by 2030 to be on a net-zero pathway (USD 700 billion).

While the positive effects of a more hydrogen-based economy are relatively established (e.g., lower CO_2 emissions, decreased urban pollution, etc.), considerable uncertainty still surrounds the consequences of hydrogen emissions to the atmosphere, because of potential indirect GHG effects^{14,19}. Here we have focused on the impact of a more hydrogen-based energy system on tropospheric methane, the second most important greenhouse gas.

We have shown how the replacement of fossil fuel energy with green or blue hydrogen could have very different consequences for tropospheric CH₄, depending on the amount of hydrogen lost to the atmosphere and the methane emissions associated with hydrogen production (Figs. 2 and 3). Specifically, tropospheric CH_4 would decrease due to the fossil fuel displacement only if the rate of H_2 losses is kept below the critical HEI.

This is around $9 \pm 3\%$ for green H₂ (Fig. 4). The same critical value would apply to other H₂ colors that do not entail the use of fossil fuels, like white or orange H₂ extracted from underground deposits^{12,47}. The critical HEI for blue H₂ is much lower due to the CH₄ emission associated with blue H₂ production. We have found that the methane emissions in a blue H₂ economy could be higher than in a fossil fuel economy if the methane supply chain had an average leak rate above 1%. Furthermore, the superimposition of CH₄ and H₂ emissions may have undesired consequences for the tropospheric burden of CH₄. This may be a potential problem in the near term, given that steam methane reforming will be used to bridge the gap between increasing H₂ demand and limited green H₂ production capacities². Our results suggest that including hydrogen emissions would aggravate the greenhouse gas footprint of blue H₂³⁶.

In addition to the CH_4 feedback, H_2 emissions are also expected to impact ozone (O₃) and stratospheric water vapor (H₂O), with negative consequences for both air quality and radiative forcing. Accounting for these effects, we can provide a comparison between the radiative forcing of hydrogen-based and fossil fuel-based energy systems. Because both H₂ and CH₄ are short-lived gas compared to CO₂, the time horizon for this comparison is crucial⁴⁸. Here we consider 20-year and 100-year time horizons. The GWP of H_2 is estimated at 11 ± 5 (100year) and 33^{+11}_{-13} (20-year)¹⁶. The GWP of CH₄ is estimated at 28 (100year) and 80 (20-year)³⁵. In an envisioned hydrogen economy that replaces the current fossil fuel industry, the H₂ emissions could be in the range 23 to 370 Tg H_2 yr⁻¹, for a H_2 emission intensity going from 1 to 10% (Fig. 2a). These emissions would have a radiative forcing impact of 0.7-12% (100-year) and 2-35% (20-year) of the current CO₂ emissions from fossil fuels (~35 Pg CO₂ yr⁻¹). If the global H₂ economy relied on blue H₂ with a 2% methane leakage rate, methane emissions would cause an additional radiative forcing impact that is around 10% (100year) and 27% (20-year) of the current CO₂ emissions from fossil fuels. Hence, in the worst scenario, up to 22% of the climate benefits of the hydrogen economy could be offset by gas losses over a 100-year horizon. The percentage could be as large as 65% over a 20-year horizon. These values could be higher on a regional scale if the leak rate of the natural gas supply chain is above 2%.

To maximize the climate benefit of hydrogen adoption, minimizing both H_2 and CH_4 losses across the supply chain of hydrogen production will need to be a priority. On the methane side, some governments and companies have already committed to reducing the leaks from the oil and gas sector, because this could be the most costeffective and impactful action for near-term climate mitigation²¹. The International Energy Agency (IEA) estimates that, with the recent rise in natural gas prices, the abatement of methane emissions from the global gas and oil sector could be implemented at no net cost⁴⁹. Hence, the accomplishment of this mitigation is only a matter of political will for the limited number of companies involved.

On the hydrogen side, the global value chain still has to be built. This offers the advantage of tackling the hydrogen emission problem ahead of time. On the one hand, energy companies will have a great interest in minimizing economic loss and safety risks due to hydrogen leaks. On the other hand, however, many technological challenges still need to be addressed. First, H₂ containment may remain an issue even as technologies progress. The high diffusivity of the small H₂ molecule has already challenged the scientific community's ability to measure the H₂ concentration in the atmosphere⁵⁰ and in the firn air of ice sheets⁵¹. Second, while more field-based estimates of H₂ losses are needed, there is currently no commercially available sensing technology able to detect small H₂ leaks at the ppb level⁴⁸. Third, global-space monitoring, which is bringing a much-needed transparency to the

quantification of real methane emissions^{27,31}, will also require new technology since H_2 , unlike CH_4 or CO_2 , does not absorb infrared radiation. For all these reasons, the uncertainty about future emissions from the H_2 value chain remains large.

Our versatile atmospheric model allowed a broad exploration of scenarios in a hydrogen-based energy system. Simulations with high resolution three-dimensional atmospheric chemistry models, which are more comprehensive but more computationally demanding, could refine our results for specific scenarios. In particular, a more detailed model could improve the assessment of H₂ displacement of fossil fuels by accounting for the emission changes of other chemical species, like CO and NO_x, which impact the CH₄–H₂–OH dynamics. Further analyses could also refine the potential changes in emission inventories due to H₂ displacement of different fossil fuels.

Methods

The Model

With the increasing anthropogenic alteration of atmospheric chemistry, detailed three-dimensional atmospheric chemistry models have become critical to evaluate the atmospheric interactions with the climate forcing^{52,53}. Nonetheless, thanks to their versatility, simplified models of atmospheric chemistry have also proven very useful to investigate the fundamental processes governing the coupling between atmospheric gases and the consequences of their possible perturbations (e.g., refs. 54–59). The insights obtained with the CH₄–CO–OH model by Prather et al.⁵⁴, in particular, led to a +40% revision of the IPCC's GWP for CH₄⁶⁰. Here we extend Prather's seminal model by adding the mass balance equation for atmospheric H₂. The purpose is to identify the key components that control the H₂ feedback on the tropospheric dynamics of CH₄ (Fig. 1).

The chemical reactions considered are

$$CH_4 + OH \xrightarrow{k_1} \dots \longrightarrow \alpha H_2 + CO \dots, R_{CH_4} = k_1[OH][CH_4],$$
⁽¹⁾

$$H_2 + OH \xrightarrow{k_2} \dots, R_{H_2} = k_2[OH][H_2],$$
 (2)

$$\text{CO} + \text{OH} \xrightarrow{k_3} \dots, \quad R_{\text{CO}} = k_3[\text{OH}][\text{CO}], \quad (3)$$

$$X + OH \xrightarrow{k_4} \dots, R_X = k_4[OH][X],$$
 (4)

with *R* representing the rates of reactions, $[\cdot]$ the concentrations, and k_i the rate coefficients. We indicated only the products with which we are concerned, the CO and H₂ produced by oxidation of CH₄(1). H₂

production through CH₄ oxidation has yield $\alpha \approx 0.37^{13}$. X encompasses all the other species, besides CH₄, CO, and H₂, that consume OH. Based on the above reactions, the balance equations for the CH₄-H₂-CO-OH system are

$$\frac{d[CH_4]}{dt} = S_{CH_4} - R_{CH_4} - R_{s'}$$
(5)

$$\frac{d[H_2]}{dt} = S_{H_2} + \alpha R_{CH_4} - R_{H_2} - R_d, \tag{6}$$

$$\frac{d[\rm CO]}{dt} = S_{\rm CO} + R_{\rm CH_4} - R_{\rm CO},$$
(7)

$$\frac{d[OH]}{dt} = S_{OH} - R_{CH_4} - R_{H_2} - R_{CO} - R_X,$$
(8)

where $R_d = k_d[H_2]$ is the H₂ uptake by soil bacteria, which plays a crucial role in the global balance of H₂ since it accounts for around 70–80% of tropospheric removal^{11,43,61}; $R_s = k_s[CH_4]$ accounts for the smaller sinks of CH₄, namely soil uptake, stratospheric loss and reactions with chlorine radicals⁶². For simplicity, we neglect the smaller sinks of H₂, i.e., stratospheric loss (~1% of removal⁶³), and CO, i.e., soil uptake and stratospheric loss (<10% of removal⁶⁴).

The solution at quasi steady state (i.e., $d[\cdot]/dt = 0$) provides the sources for fixed tropospheric concentrations. Positive solutions for OH occurs if $S_{OH} > (2 + \alpha)(S_{CH_4} - R_s) + S_{CO} + S_{H_2} - R_d$, i.e., when there is enough OH to oxidize all CO sources, the part of CH₄ sources that is not balanced by smaller sinks, and the part of H₂ sources that is not balanced by the soil uptake. The excess of OH consumed by other gases, besides CH₄, CO, and H₂, can be defined as $E_{OH} = R_X/(R_{CH_4} + R_{CO} + R_{H_2})$. The values representing average tropospheric conditions are summarized in Table 1. The values of S_{OH} and S_{CO} are kept constant in all scenarios.

Linear stability and transient dynamics

We investigate the effects of an emission pulse of H₂ on the tropospheric system (5)–(8). The timescales and modes of the atmospheric response to chemical perturbations are defined by the eigenvalues and eigenvectors of the system^{54,55}. Indicating with $\mathbf{c}(t)$ the solution vector of the system (5)–(8), the temporal dynamics of a small perturbation $\hat{\mathbf{c}}$ around \mathbf{c} evolves as

$$\frac{\mathrm{d}\hat{\mathbf{c}}}{\mathrm{d}t} = \mathbf{J}\hat{\mathbf{c}},\tag{9}$$

|--|

		CH ₄	H ₂	СО	ОН	$-\lambda_i^{-1}$ (yr)
Steady state	Concentration (ppb)	1890	530	80	10 ⁶ cm ⁻³	
	Sources (ppb/yr)	226	265ª	480ª	1333	
	τ (yr)	8.3	2	0.17	1 s	
Linear stability	CH ₄ mode	1%	0.31%	0.64%	-0.39%	12.3
	H ₂ mode	-0.01%	1%	0.03%	-0.06%	2
	CO mode	-0.008%	0.001%	1%	-0.36%	0.2
	OH mode	^b	^b	^b	1%	1.5 s

^aSources for CO and H₂ include production from CH₄ oxidation.

^b... is <10⁻⁷.

Sources are obtained from the system (5)–(8) at steady state with the current tropospheric concentrations. τ is the average lifetime of each gas. The modes are expressed as relative changes normalized so that the dominant species' ratio is 1%. Reaction rates are defined as follows: $k_1 = 3.17 \times 10^{-15} \text{ cm}^3/\text{s}$; $k_2 = 3.8 \times 10^{-15} \text{ cm}^3/\text{s}$; $k_3 = 1.9 \times 10^{-13} \text{ cm}^3/\text{s}$; $k_8 = 0.02 \text{ yr}^{-1}$; $k_d = 0.38 \text{ yr}^{-1}$ is such that soil uptake accounts for 75% of atmospheric H₂ removal; $k_4[X] = 0.38 \text{ s}^{-1}$ ($E_{OH} = 0.82$) is defined so that 45% of OH is consumed by the species X, 36% by CO, 14% by CH₄, and 5% by H₂³⁸. Concentrations are converted to mixing ratios using 1 ppb = 1.57 \times 10^{10} \text{ cm}^{-3}; sources are converted from ppb/yr to Tg/yr using 4.22 \times 10^{18} \text{ kg} as the troposphere mass⁶⁸.







Colors highlight the contributions of the different modes. When different modes superimpose, the faster-decaying mode is shown on top of the others.

where **J** is the Jacobian of the system evaluated in **c**. For the equilibrium solution \mathbf{c}_0 representing the current tropospheric concentrations, the eigenvalues and eigenvectors, or modes, of the linearized system (9) are reported in Table 1. Since all eigenvalues are real and negative ($\lambda_i < 0$), the equilibrium solution \mathbf{c}_0 is a stable node. As a result, any small perturbation asymptotically decays in time with a timescale defined by the negative reciprocal of the eigenvalue.

Because the system equations are coupled, the decay timescale $(-\lambda_i^{-1})$ of a gas perturbation does not necessarily correspond to the gas steady state average lifetime (τ_i) . The CH₄ perturbation, in particular, decays with a timescale that is much larger than what predicted by its steady state lifetime, i.e., $R = -\lambda_{CH_4}^{-1}/\tau_{CH_4} > 1$. This mechanism, known as the CH₄ feedback effect^{55,65}, has a crucial role in increasing the GWP and the environmental impact of CH₄ emissions. Detailed models of atmospheric chemistry usually provide *R* around 1.3–1.4⁶⁵. We find a marginally higher feedback factor, namely $R \approx 1.5$, in agreement with previous findings using Prather's box model^{54,55,57}. The decay timescale of the H₂ perturbation instead corresponds to the H₂ average lifetime, namely $-\lambda_{H_2}^{-1} \approx \tau_{H_2}$, in agreement with results from detailed atmospheric chemistry model⁵¹.

While the modal eigenvalue analysis correctly captures the asymptotic stability of the solution \mathbf{c}_{0} , it does not describe the perturbation dynamics at finite times, i.e., before the asymptotic decay. Still within the domain of the linearized system (9), a more complete picture can be obtained by analyzing the temporal evolution of the solutions with specific attention to the emergence of transient growth phenomena, which are known to occur in systems where the modes are non-orthogonal, as in the present case. When large enough, a transient growth can even trigger nonlinearities that destabilize the equilibrium solution⁶⁶.

Figure 5 shows the transient growth phase of tropospheric CH₄ and CO that follows a 10% perturbation of H₂ concentration. Specifically, the pulse of H₂ causes a drop in OH and a build-up of CH₄ that lasts a few years, while the H₂ perturbation decays with the timescale τ_{H_2} . The CH₄ build-up then decays in the same manner as would a direct pulse of CH₄ with a timescale defined by the CH₄ feedback effect. In analytical terms, the perturbation of tropospheric CH₄ mainly due to the excitation of H₂ and CH₄ modes is given by δ [CH₄] \approx 2.76 $e^{\lambda_{CH_4}t}$ - 2.82 $e^{\lambda_{H_2}t}$ + 0.06 $e^{\lambda_{CO}t}$.

Using this result in traditional GWP formulas³⁵ yields a GWP for H₂ due to direct CH₄ perturbation around 7.8 with the 100-year timehorizon and 22 with the 20-year time horizon. It is estimated that around half of the H₂ indirect radiative forcing is due to the direct CH₄ perturbation, and the other half to the O₃ and stratospheric H₂O impacts caused by both H₂ and H₂-induced CH₄ perturbations¹⁴. Taking this into account yields a total GWP for H₂ of 15.6 with the 100-year time-horizon and 44 with the 20-year time-horizon. These values are in the upper range of the recent estimates of 11 ± 5 for GWP100 and 33⁺¹¹₋₁₃ for GWP20 obtained with a detailed model of atmospheric chemistry¹⁶. Notably, the consequences of the H₂ pulse on CH₄ are relatively small in magnitude because most of the additional H₂ is oxidized by soil bacteria and not by OH. The stability of this biotic sink as affected by climate change and anthropic pressure is hence a crucial aspect for the impact of future H₂ emissions, as further discussed in the main text.

Critical hydrogen emission intensity

We here derive an explicit expression for the critical H₂ emission intensity (HEI_{cr}) for methane mitigation, defined as the emission rate that offsets the H₂ replacement of fossil fuels. The expression is derived for an infinitesimal replacement of fossil fuel energy with H₂ (d*E* in ExJ/yr), but well approximates the critical HEI for finite replacement of fossil fuel energy (see Supplementary Fig. 3). As a first step, we differentiate the system (5)–(8) at equilibrium (d[·]/d*t* = 0) with respect to *E*. This yields

$$S_{CH_4,E} - k_1[CH_4][OH]_E = 0,$$
 (10)

$$S_{H_2,E} + \alpha k_1 [CH_4] [OH]_E - k_2 ([H_2][OH])_E - k_d [H_2]_E = 0, \quad (11)$$

$$k_1[CH_4][OH]_E - k_3([CO][OH])_E = 0,$$
 (12)

 $k_1[CH_4][OH]_E + k_2([H_2][OH])_E + k_3([CO][OH])_E + k_4[X][OH]_E = 0,$ (13)

where subscript *E* indicates $d \cdot /dE$. [CH₄]_{*E*} = 0 because of the definition of the critical H₂ emission intensity, which leaves the methane concentration unaltered. We consider that only H₂ and CH₄ sources vary with *E*, while $S_{OH,E} = S_{CO,E} = 0$. These variations can be estimated as

$$S_{H_2,E} = a_{H_2} \left(-ff_{H_2} + \frac{HEI}{\eta_{H_2}(1 - HEI)} \right),$$
 (14)

$$S_{CH_{4},E} = a_{CH_{4}} \left(-ff_{CH_{4}} + \frac{r \text{ MEI}}{\eta_{H_{2}}(1 - \text{HEI})} \right),$$
 (15)

where HEI and MEI are the hydrogen and methane emission intensities, respectively (MEI = 0 for green H₂); η_{H_2} is H₂ higher heating value; *r* is the amount of CH₄ needed to produce a unit of blue H₂; ff_{CH₄} and ff_{H₂} are the average amounts of CH₄ and H₂ emitted per ExJ of fossil fuel energy; a_{H_2} and a_{CH_4} are conversion factors.

Substituting Eqs. (14), (15) into the system (10)–(13) and after some algebra, one obtains the critical H_2 emission intensity

$$\text{HEI}_{\text{cr}} = \frac{A\left(\text{ff}_{\text{CH}_4} \eta_{\text{H}_2} - r \text{ MEI}\right) + B \text{ ff}_{\text{H}_2} \eta_{\text{H}_2}}{A \text{ ff}_{\text{CH}_4} \eta_{\text{H}_2} + B\left(\text{ff}_{\text{H}_2} \eta_{\text{H}_2} + 1\right)}.$$
 (16)

where the dependence to the atmospheric composition is embedded in $A = k_d(k_4[X] + k_2[H_2] + 2k_1[CH_4]) + k_2[OH]((\alpha + 2)k_1[CH_4] + k_4[X])$ and $B = 8k_1k_2[CH_4][OH]$. Parameters have been defined as follows: $\eta_{H_2} = 0.143 \text{ ExJ/Tg}_{H_2}$, $r = 3.2 \text{ kg}_{CH_4}/\text{kg}_{H_2}$, $ff_{CH_4} = 0.225 \text{ Tg}_{CH_4}/\text{ExJ}$, $ff_{H_2} = 0.0225 \text{ Tg}_{H_2}/\text{ExJ}$, $a_{CH_4} = 0.43 \text{ ppb/Tg}$, $a_{H_2} = 8a_{CH_4}$. To obtain the value of r, we used the estimate of 3.7 kg of natural gas for kg of H2⁶⁷, which includes feedstock and energy requirements, and we assumed that 85% of natural gas by weight is composed by methane. ff_{CH_4} and ff_{H_2} are obtained as the ratio between the global CH₄ and H₂ emissions due to fossil fuel use and the global fossil fuel energy.

Data availability

All data generated during this study are provided in the supplementary dataset file.

Code availability

The code used to generate the results is provided in the supplementary dataset file.

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Author contributions

M.B.B., S.W.P., and A.P. conceptualized the work. M.B.B. developed the analytical model with contributions from F.P., analyzed the results and prepared the manuscript. A.P., F.P., and S.W.P. supervised the work and edited the manuscript.

Competing interests

The authors declare no competing interests.

Additional information

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