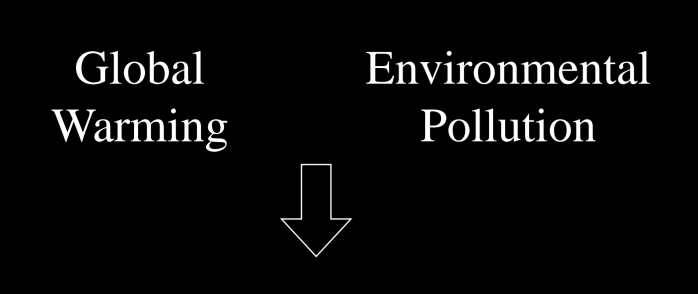
## MXenes as Photocatalytic Materials for Water Splitting



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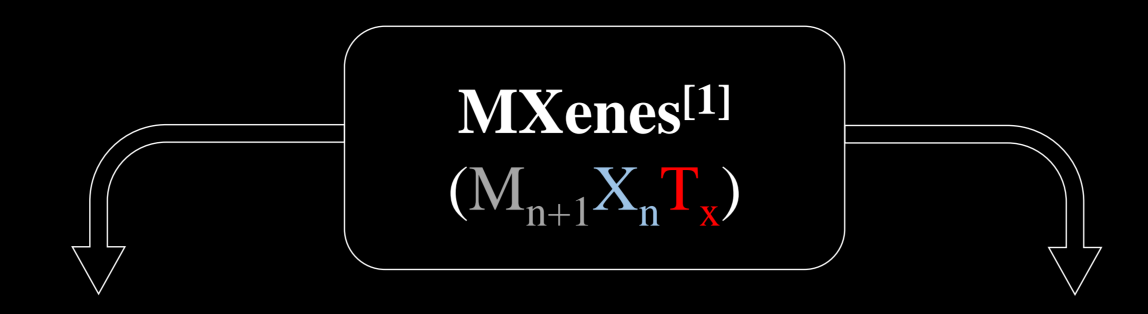


Need for green fuels

Hydrogen (H<sub>2</sub>)



Modify and design the bandgap of by varying



**CBM** 

**VBM** 

M = Transition Metal (Groups III - VI) $X = C \text{ or } N \quad n = 1-3$ T = Termination (p-block: O, F, OH, H, S, Cl)

 $H_2$  %

Semiconductors when adding a termination.<sup>[2]</sup> Good candidates for photocatalysing the water splitting process and produce clean H<sub>2</sub>.<sup>[3]</sup>

 $H^+/H_2$ 

 $O_2/H_2O$ 

Sustainable source of H<sub>2</sub>: Water Splitting  $2 H_2O \rightarrow 2 H_2 + O_2$ 

#### **Problem:** Requires larges amounts of energy

**Solution:** Sunlight as energy source

#### Photocatalysis



## TOOLS \*

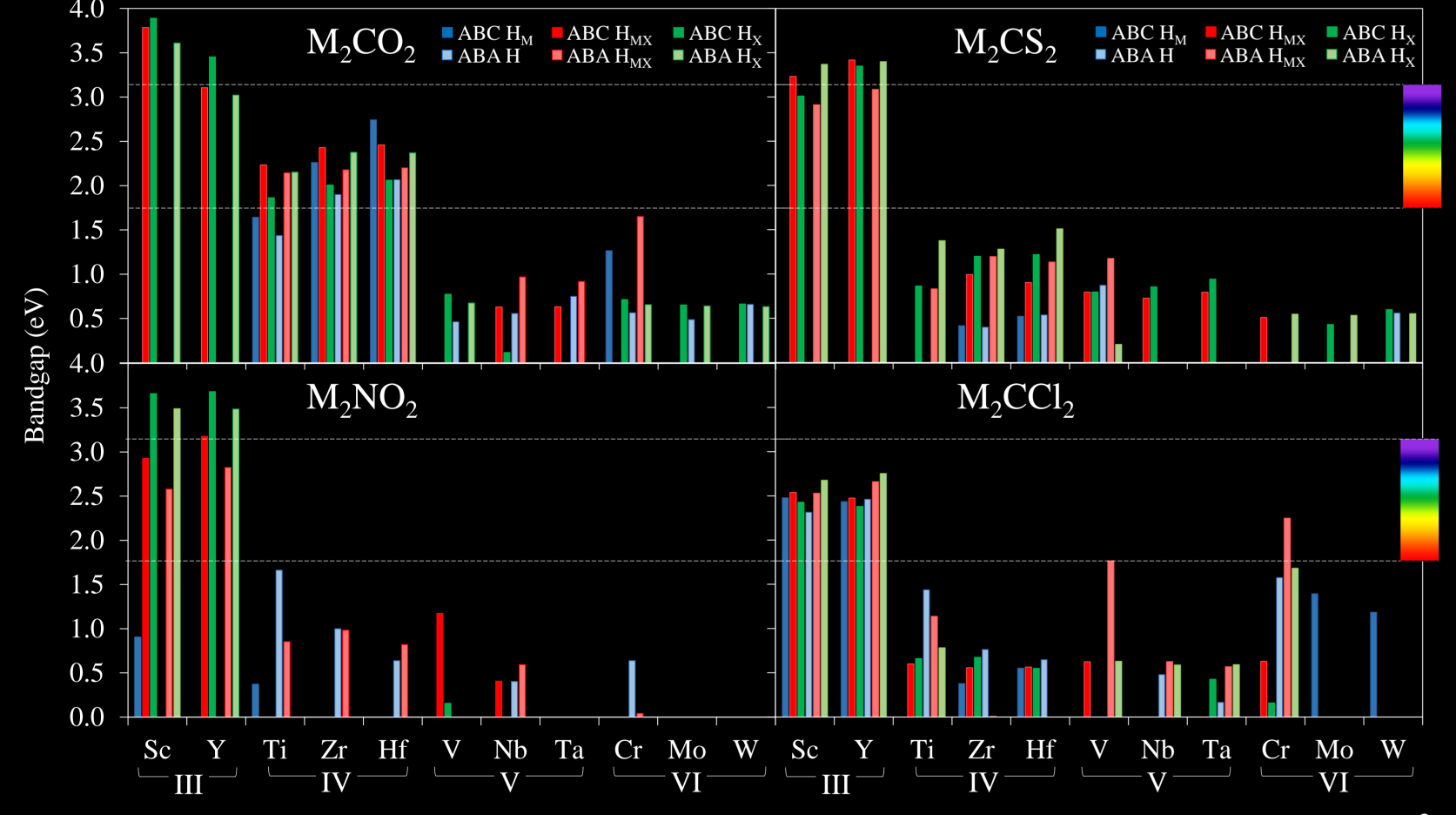
**Computational Method: DFT** Functional: PBE and PBE0 Structures: periodic slab models, considering two stackings (ABC and ABA) and three termination positions for each stacking (H<sub>M</sub>/H,

 $H_{MX}$  and  $H_{X}$ )

# OBJECTIVES Q

**MXenes** their composition (M, X, T), width (n), stacking and termination position, to find potential photoactive candidates for water splitting using solar light.

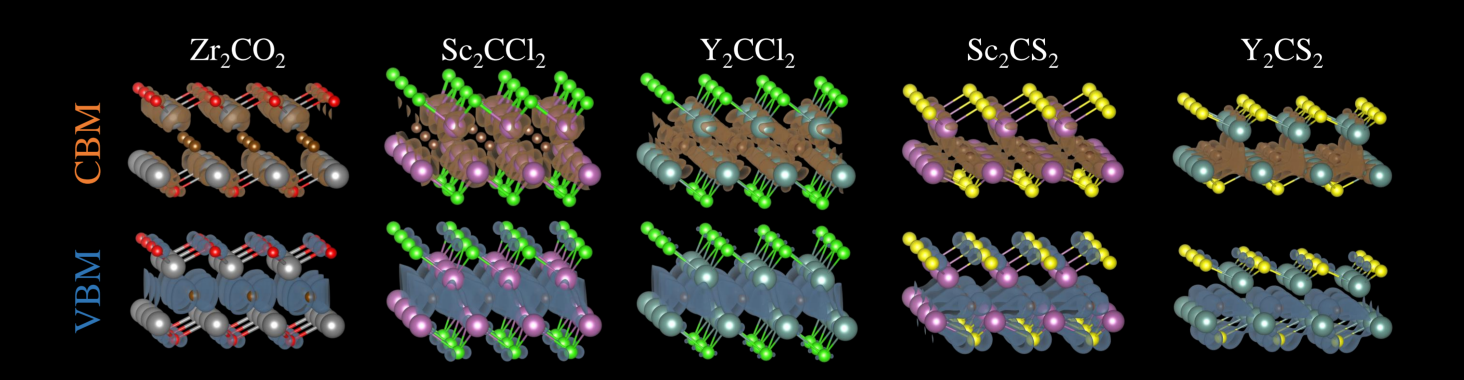
## Density of States



- 6 different structures for each terminated MXene are considered (2376 in total).
- Group III and IV MXenes with  $n = 1 \rightarrow large bandgaps$ , in the visible region. The most promising cases for being photoactive materials with sunlight.
- C-MXenes → more semiconducting cases and larger bandgaps than N-MXenes.
- Pristine MXenes  $\rightarrow$  metallic properties (not photoactive).

 $H_2O$ 

MXenes  $n = 2, 3 \rightarrow$  Increasing the amount of "bulk" tends to make them metallic.



## X Band Alignment

- The band alignment with respect to the half-reaction potentials has been studied for the most promising photoactive cases ( $E_g > 1.23 \text{ eV}$ ).
- The ideal cases will be those that, in addition to having a suitable band alignment, are the most stable structure among the six considered.
- Several structures from Groups III and IV exhibit correct alignments.
- The cases of Zr<sub>2</sub>CO<sub>2</sub>, Sc<sub>2</sub>CCl<sub>2</sub>, Y<sub>2</sub>CCl<sub>2</sub>, Sc<sub>2</sub>CS<sub>2</sub>, and Y<sub>2</sub>CS<sub>2</sub> fulfill these optimal conditions, which allows us to propose them as potential candidates for the photocatalytic water splitting.
- These systems exhibit an indirect bandgap and a significant charge density separation between the valence band maximum and conduction band minimum (VBM and CBM), promoting the separation of generated charges.

#### Conduction Band Bandgap 2.26 eV 2.48 eV 2.44 eV 3.23 eV 3.42 eV $H_2O$ Valence Band $Zr_2CO_2$ Sc<sub>2</sub>CCl<sub>2</sub> Y<sub>2</sub>CCl<sub>2</sub> Sc<sub>2</sub>CS<sub>2</sub> $Y_2CS_2$

## CONCLUSIONS \

Based on DFT calculations, MXenes with n = 1, X = C, M = Groups III and IV,and T = O, S, Cl, especially  $Zr_2CO_2$ ,  $Sc_2CCl_2$ ,  $Y_2CCl_2$ ,  $Sc_2CS_2$ , and  $Y_2CS_2$ , exhibit a visible range bandgap, optimal for solar light absorption, and band edges that exceed the half-reaction potentials of water splitting, suitable for photocatalysing the process and generating green H<sub>2</sub>.

#### REFERENCES



[2] Adv. Funct. Mater. 2013, 23, 2185–2192. [3] J. Mater. Chem. A 2016, 4, 11446–11452.



