Supporting Information

Metallic Tungsten Carbide Nanoparticles as Near-infrared Driven Photocatalyst

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Figure S1 BET and corresponding pore size distribution curves.

Figure S2 W 4f XPS spectra of WC nanoparticle before and after Ar$^+$ etching 20 nm.

Table S1 Atomic analysis of W 4f before and after Ar$^+$ etching

<table>
<thead>
<tr>
<th>Chemical state</th>
<th>Atomic ratio</th>
<th>Before etching</th>
<th>After etching</th>
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<tbody>
<tr>
<td>W$^{4+}$(W-C)</td>
<td>0.55</td>
<td>0.73</td>
<td></td>
</tr>
<tr>
<td>W$^{5+}$(W-O)</td>
<td>0.15</td>
<td>0.19</td>
<td></td>
</tr>
<tr>
<td>W$^{6+}$(W-O)</td>
<td>0.30</td>
<td>0.08</td>
<td></td>
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</table>
Figure S3 Temperature profiles of (a-d) the 1 mg/L WC aqueous dispersion without cooling water, (e-h) water without cooling water, (i-l) the 1 mg/L WC aqueous dispersion with cooling water under 30 min NIR irradiation.

Figure S4 (a) Temperature-radiation time dependent of different solution. (b) $\frac{C_t}{C_0}$ of 10 mg/L MB in 85 °C water bath (under dark condition).

Figure S5 DFT calculated density of state (DOS) and the corresponding projected DOS of WC.

The DFT calculation of electronic structure of WC was performed on CATSEEP, which is one of functional module in Material Studio package. The gradient corrected exchange-correlation with Perdew-Burke-Ernzerhof functional (GGA-PBE) and ultrasoft pseudopotential plane-wave methods were used for all the calculations. A hexagonal 4x4x2 supercell with 64 atoms was built for simulating WC. The cut-energy
of 340 eV and homogenate meshed 3x3x3 k-point were used throughout the calculation. The self-consistent geometry optimization was performed until it satisfies the convergence tolerances (0.01 GPa and 5x10-6 eV for maximum stress and energy change).

Figure S6 NIR photocatalytic degradation on Phenol of WC nanoparticles.

Figure S7 Recycle NIR photocatalytic degradation of MB for 4 runs.

Figure S8 XRD spectra of WC nanoparticles before and after 4 run’s photocatalyst.