The Acetone synthesis

- Cumene hydroperoxide decomposition CC(C)(00)c1cccc1>acid>CC(C)=0
- 2. Isopropanol dehydrogenation CC(C)0>Cu>CC(C)=0
- 3. Propene hydration/oxidation {2}C=CC.O=O>Pd/Cu>{2}CC(C)=O

```
from rxnSMILES4AtomEco import atom_economy

# calculate Verbose mode
atom_economy("{2}C=CC.0=0>Pd/Cu>{2}CC(C)=0")
```

Atom Economy Calculation: REACTANTS SMILES: C=CC Molecular Formula: C3H6 Molecular Weight: 42.08 g/mol Coefficient: 2.0 SMILES: 0=0 Molecular Formula: 02 Molecular Weight: 32.00 g/mol Coefficient: PRODUCTS SMILES: CC(C)=0

Molecular Formula: C3H6O

Molecular Weight: 58.08 g/mol

Coefficient: 2.0

Atom Economy: 100.0%

```
reactions_smiles_pathway = {
    # Cumene hydroperoxide decomposition
    "CC(C)(00)c1ccccc1>acid>CC(C)=0": "Cumene",
    # Isopropanol dehydrogenation
    "CC(C)0>Cu>CC(C)=0": "Isopropanol",
    # Propene oxidation
    "{2}C=CC.0=0>Pd/Cu>{2}CC(C)=0": "Propene"
}
```

```
from rxnSMILES4AtomEco import get_atom_economy

# Store results
results_paths = {}

# Loop through pathways and calculate atom economy
for smiles, name in reactions_smiles_pathway.items():
    results_paths[name] = get_atom_economy(smiles)

# Print results
for pathway, value in results_paths.items():
    print(f"{value:.1f}% {pathway}")
```

```
38.2% Cumene
96.6% Isopropanol
100.0% Propene
```

```
import matplotlib.pyplot as plt

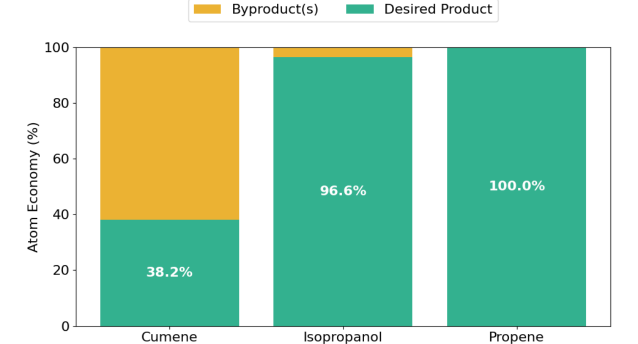
# Extract labels (pathway names) and values (atom economy)

# Labels for x-axis
pathways = list(results_paths.keys())

# Y-axis values
atom_economy_values = list(results_paths.values())
```

```
# Calculate byproducts as (100 - atom economy)
byproduct_values = [100 - value for value in atom_economy_values]
# Create a single plot
# Single axis for both plots
fig, ax = plt.subplots(figsize=(10, 7))
plt.rcParams.update({'font.size': 16})
# Plot bars: green for desired product, orange for byproducts
bars = ax.bar(pathways, atom_economy_values, color='#009E73', \
              label="Desired Product", alpha=0.8)
ax.bar(pathways, byproduct_values, bottom=atom_economy_values, \
       color='#E69F00', label="Byproduct(s)", alpha=0.8)
# Annotate bars with atom economy percentages
for bar in bars:
   height = bar.get_height()
    ax.text(bar.get_x() + bar.get_width()/2, height/2, f'{height:.1f}%',
            ha='center', va='center', color='white', fontweight='bold')
# Titles & labels
ax.set_title("Acetone Atom Economy Across Pathways\n\n")
ax.set_xlabel("\nSynthesis Pathways")
ax.set_ylabel("Atom Economy (%)")
# Limit y-axis to 100%
ax.set_ylim(0, 100)
ax.legend(loc='upper center', bbox_to_anchor=(0.5, 1.2), ncol=2,reverse=True)
# Improve layout
plt.tight_layout()
# Show the plot
plt.show()
# To save the plot, if used, comment line above and uncomment lines below
#save_path = './acetone_histogram.png' # Writable on many systems
#plt.draw() # Force render
#print(f"Saving to: {save_path}")
#plt.savefig(save_path, dpi=300, bbox_inches='tight')
#plt.close()
#print("Save completed (check directory).")
```

Acetone Atom Economy Across Pathways



Synthesis Pathways

```
# Function to clean and expand reaction SMILES
def process_smiles(smiles):
    # Remove reagents/catalysts (text between ">...>")
    cleaned_smiles = re.sub(r'>[^>]+>', '>>', smiles)
    # Expand {n} notation: e.g. "{2}C=CC" \rightarrow "C=CC.C=CC"
    def expand match(match):
       # Extract number
       n = int(match.group(1))
        # Extract SMILES
        molecule = match.group(2)
        # If stoichiometric coefficient found, repeat SMILES n times
        # (works only with integer coefficients!)
        return ".".join([molecule] * n)
    expanded smiles = re.sub(r'(d+))([A-Za-z0-9@+-=#()[]]+)', 
                             expand_match, cleaned_smiles)
    return expanded_smiles
# Loop through each reaction and plot separately
for reaction_smiles, name in reactions_smiles_pathway.items():
    # Process SMILES
    cleaned_smiles = process_smiles(reaction_smiles)
    print(f"Processing: {name}, SMILES: {cleaned_smiles}")
    try:
        # Convert to RDKit reaction object
        reaction = rdChemReactions.ReactionFromSmarts(cleaned smiles, \
                                                      useSmiles=True)
        # Generate reaction image
        img = Draw.ReactionToImage(reaction, subImgSize=(300, 300))
        # Save path
        save_path = os.path.join(output_dir, f"{name}.png")
        img.save(save_path, format="PNG", dpi=(300, 300))
        print(f"Saved: {save_path}")
    except Exception as e:
        print(f"Error processing {name}: {e}")
```

else:

print("Script execution skipped.")

Script execution skipped.