Air Pollution and SI Engine Emissions

Atmospheric Pollution

• <u>SMOG</u>

O₃ NO₂
– Ozone Nitrogen dioxide

R-C-OONO₂
PAN(Peroxyacyl Nitrate)

TOXICS

 CO, Benzene, 1-3 butadiene, POM (Polycyclic organic Matters), Aldehydes

Primary Pollutants: Direct emissions from vehicles > CO, HC, NOx, PM(Particulate matters), SOx, aldehydes

Secondary Pollutants: From interaction of emissions with the atmosphere

> O₃, PAN, NO₂, Aldehydes

Atmospheric Pollution

Smog formation:

Acid rain:

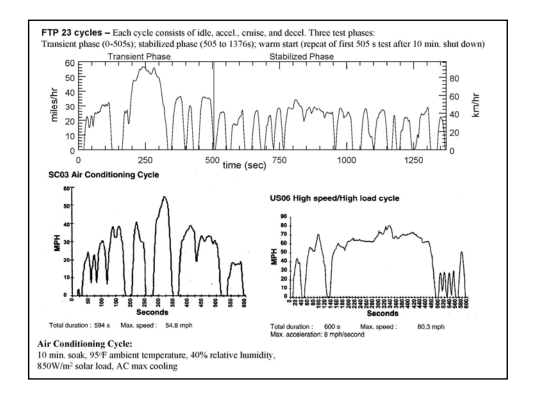
$$SO_{2} + \dot{O}H \rightarrow H\dot{O}SO_{2}$$

$$H\dot{O}SO_{2} + O_{2} \rightarrow H\dot{O}_{2} + SO_{3}$$

$$SO_{3} + H_{2}O \rightarrow H_{2}SO_{4}$$

$$NO + \frac{1}{2}O_{2} \rightarrow NO_{2}$$

 $NO_2 + \dot{O}H \rightarrow HNO_3$



EMISSIONS MECHANISMS

- · CO emission
 - Incomplete oxidation of fuel under fuel rich conditions
- NOx emisison
 - Reaction of nitrogen and oxygen in the high temperature burned gas regions
- Particulate matter (PM) emission (most significant in diesel engines; there are significant PM emissions in SI engines in terms of number density, especially in direct injection engines)
 - Particulates formed by pyrolysis of fuel molecules in the locally fuel rich region and incomplete oxidation of these particles
 - Lubrication oil contribution
- · Hydrocarbon emissions
 - Fuel hydrocarbons escape oxidation (or only partially oxidized) via various pathways

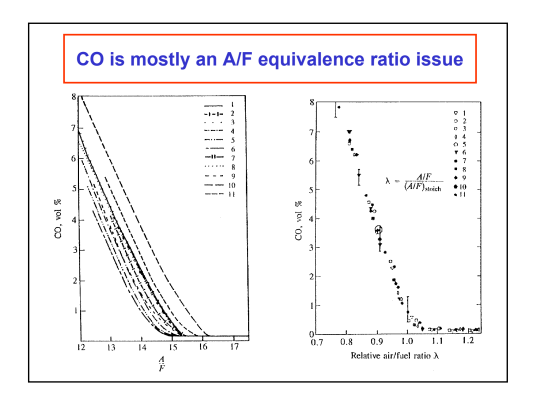
Typical steady state SI engine-out emissions

- NOx is a few thousand parts per million
- CO is around 0.5-1% for stoichiometric operation
- HC is 500-2000 ppm for fully warm up engine
- PM very small by mass

CO Emissions Mechanism

- CO is the incomplete oxidation product of the fuel carbon
- Significant amount in fuel rich condition
- Immediately following combustion, CO is in chemical equilibrium with the burned gas
- During expansion, as the burned gas temperature decreases, CO is 'frozen'
 - Empirical correlation

$$\frac{[CO][H_2O]}{[CO_2][H_2]} \approx 3.7$$



NO FORMATION CHEMISTRY

Zeldovich Mechanism

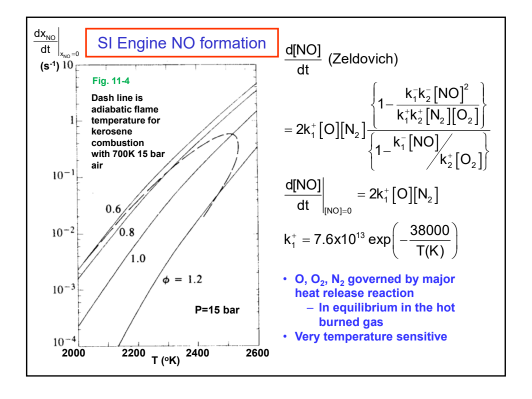
$$N_2 + O \Leftrightarrow NO + N$$
 (1)

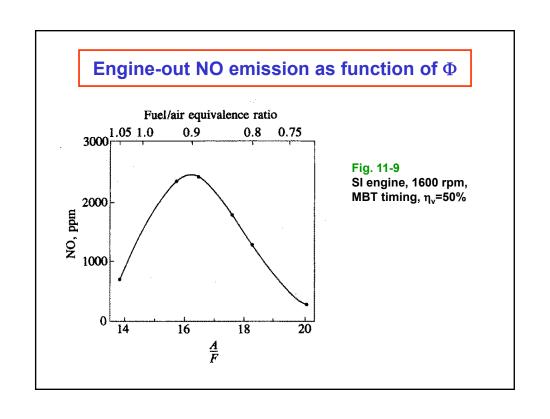
$$N + O_2 \stackrel{\kappa_2}{\longleftrightarrow} NO + O$$
 (2)

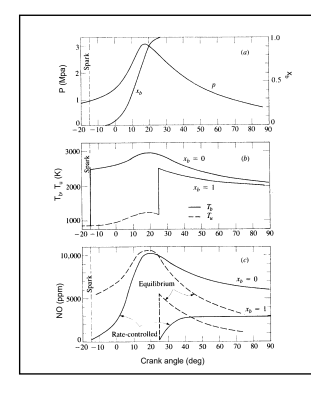
$$N + OH \stackrel{K^*_3}{\underset{k_3}{\longleftarrow}} NO + H$$
 (3) (extended Zeldovich Mechanism*)

- > NO formation is kinetically controlled
- ➤ Reactions involving N is fast; N is in steady states (d[N]/dt ≈ 0)
- ➤ Very temperature sensitive
- At high temperature (≥ 1000K), equilibrium favors NO versus NO₂ formation
 - ightharpoonup Engine-out [NO₂]/[NO_x] $\leq 2\%$

*Extended Zeldovich mechanism is quantitatively important; see text: discussion immediately following Eq. (11.7), and Table 11.2.





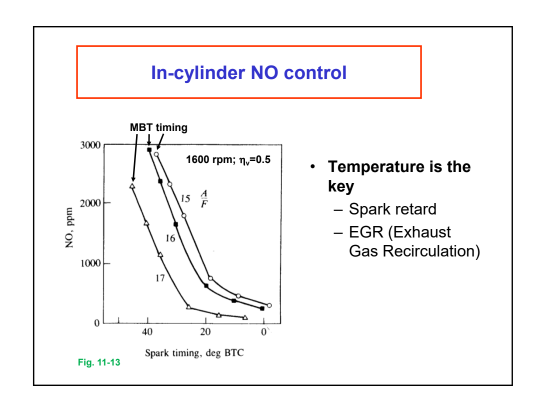


Thermodynamic state of charge

Fig. 9-5 Cylinder pressure, mass fraction burned, and gas temperatures as function of crank angle during combustion.

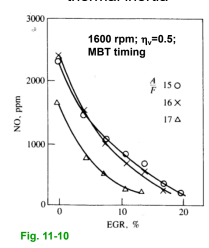
- NO formed in burned gas
- Different "layers" of burned gas have substantially different temperature, hence different amount of NO production
- In reality, there is mixing between the layers
- Rate is non-linear in temperature

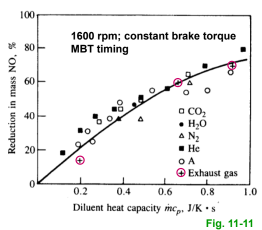
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NO control by EGR

- EGR is a dilution effect
 - Reduce burned gas temperature via increase in thermal inertia





HC emissions

- Importance
 - Photochemical smog (irritant; health effects)
 - Significant loss of fuel energy
- Measurement
 - Flame Ionization Detector (FID)
 - > Chemi-ionization process
 - > Signal proportional to C atom concentration
- Emissions regulation: NMOG as g/mile
 - EPA definition of HC
 - ➤ Normal gasoline CH_{1.85}
 - ➤ Reformulated gasoline CH_{1.92}
 - ➤ Compressed natural gas CH_{3.78}
 - Need speciation to detect CH₄

HC Impact on smog formation

- Species dependent
 - Assessed as MIR of individual VOC
- VOC = volatile organic compounds

Kinetic reactivity =
$$\frac{\text{VOC reacted}}{\text{VOC input}}$$

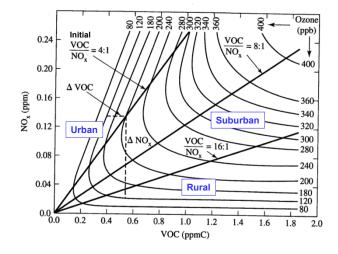
Mechanistic reactivity =
$$\frac{\text{Ozone formed}}{\text{VOC input}}$$

Maximum Incremental Reactivity (MIR)

$$\label{eq:mirror} \text{MIR} = \frac{m_{\text{ozone,test case; max}} - m_{\text{ozone,base case; max}}}{\text{VOC increment to base case}}$$

EKMA (Empirical Kinetic Modeling Approach) methodology: follow air column (Lagrangian) from 0800 using O3 as indicator. Maximum O3 formation occurs at about 1500-1700 hr.

Ozone production



EKMA plot of the maximum ozone concentration as a function of the initial Nox and VOC concentrations; from Fay & Golomb, *Energy and the Environment*, Oxford 2002.

	MIR		MIR		
l .		i .	(g O3/g NMOG)		
Cit	7 290	Benzene	0.420		
			2.730	Carter Index for	
		Ethylbenzene	2.700	Ozone Forming	
		M-Xylene	8.160	Ozone Forming	
			6.600	Potential	
			2.220		
			6.460	(CARB July, 1992)	
				(0,112 04.), 1002,	
		1.3-Methylethylbenzene			
			7.200		
		1.3.5-Trimethylbenzene	10.120		
		1.2-Methylethylbenzene			
	6.220	1,2,4-Trimethylbenzene			
	4.900	Iso-Butylbenzene			
	9.080	1,2,3-Trimethylbenzene			
	8,800	Indane	1.060		
	8.800	1,3-Diethylbenzene			
	6.410	1.4-Diethylbanzene			
	7.660	1,2-Diethylbenzene		Table from SAE	
				Table ITOIII SAE	
				Paper 932718	
		1.2 Dimethyl-2 Ethylbenzene			
		1.3 Dimethyl-2 Ethylbenzene	9.070	(Tauchida et.al)	
		1.2.4.5-Tetramethylbenzene		(,	
		1.2.3.5-Tetramethylbenzene	9.070		
			1.060		
		1.2.3.4-tetramethylbenzene	9.070		
	6.690	Methyl 1-Bulyl Ether	0.620		
	5.690	Ethyl 1-Bulyl Ether	1,980		
		Methanol			
2-Mathyl-Hexaue	6.630	Ethanol	1.340		
	5.530				
	5.520	Formaldehyde	7.150		
	5.290	Acetaldehyde	5.520		
p-4 coming		Acrolein	6.770		
		Propionaldehyde	6.530		
		Acetone	0.560		
Matha	dalamı avı	dained in CAE I	2000 000	740	
wetno	aology exp	Jiaineu in SAE i	-aper 900	J <i>I</i> 10	
		and Camen	-		
(Lowi and Carter)					
(g O3) MIR	Q OSIP (MMOG)	GO 009 NMOG GO 309 NMOG GO 309 NMOG GO 305 NMOG GO 3	10	March Graph Grap	

HC sources

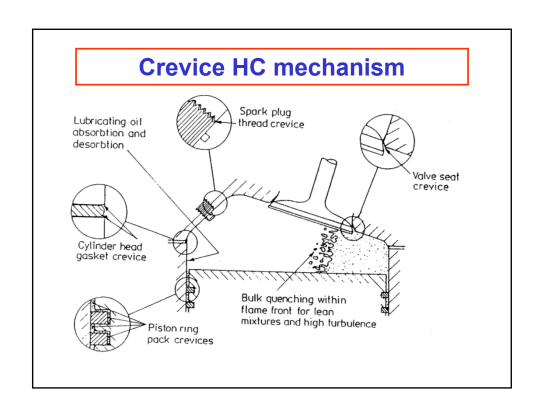
- Non-combustion sources
 - Fueling loss
 - Diurnal emissions
 - Running loss
 - Hot soak
 - Blow by
 - ➤ A few L/min; depends on load and RPM
 - \triangleright At light load, 1500 rpm, blow by \sim 4L / min

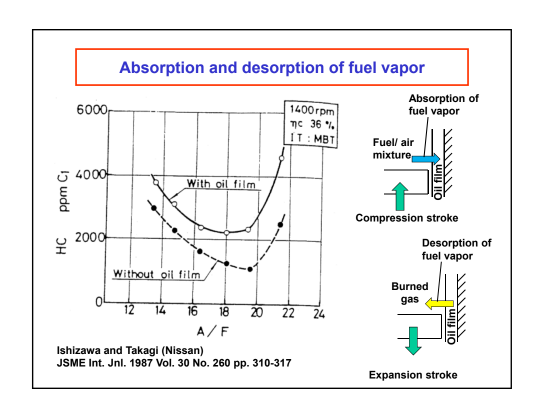
HC sources (cont.)

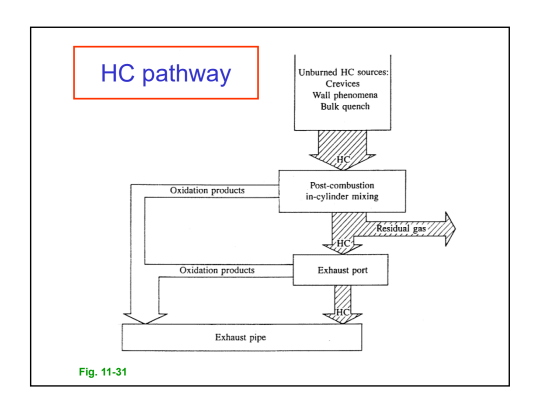
- Combustion sources
 - 300 to 3000 ppmC1 typical➤ Stoichiometric mixture is ~120,000 ppmC1
 - Main combustion: very little HC except for very lean/ dilute or very late combustion (misfires/ partial burns)
 - ➤ Various mechanisms for HC to escape from main combustion
 - Cold start emissions (wall film) especially important

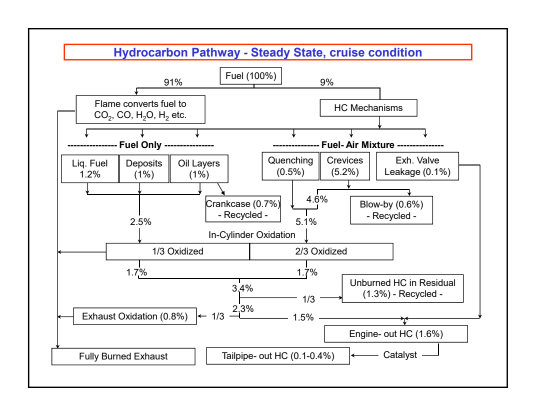
SOURCES OF UNBURNED HC IN SI ENGINE

- a) Crevices
- b) Absorption and desorption in oil layers
- c) Absorption and desorption in deposit
- d) Quenching (bulk and wall layer)
- e) Liquid fuel effects
- f) Exhaust valve leakage









HC Sources: Magnitudes and Percent of Total Engine-out Emissions*

(SAE Paper 932708)

Source	% Fuel Escaping Normal Combustion	Fraction Emitted as EOHC	% Fuel as HC Emissions	% of Total EOHC Emissions
Crevices	5.2	0.15*	0.682*	42.6
Quench	0.5	0.15	0.074	4.6
Oil Layers	1.0	0.09**	0.090**	5.6
Deposits	1.0	0.30	0.300	18.7
Liquid Fuel	1.2	0.30	0.356	22.2
Valve Leakage	0.1	1.00	0.100	6.3
Total	9.0		1.60	100

Blowby (0.6%) subtracted

*steady state cruise condition (1500 rpm, 2.8 bar NIMEP)

HC control

- Reduce crevice volume
- · Keep liner hot
- · Spark retard
 - Higher burned gas temperature in the later part of expansion stroke and higher exhaust temperature
- · Comprehensive cold start strategy
 - Retard timing, fuel rich followed by exhaust air injection

Amount to crank case (0.7%) subtracted