Final Report

Referring to
Grant # 043021, Award # FA8655-04-1-3021
Period of Performance 1 February 2004 to 31 January 2005

Title: Advanced singlet oxygen generator for a COIL

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Date of submission: 31 January 2005
12 month after grant award according to Order for Supplies or Services
**Report Date:** 14-02-2005  
**Report Type:** Final Report

**Title and Subtitle:** Advanced Singlet Oxygen Generator for a COIL

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**Sponsoring/Monitoring Agency Name(s) and Address(es):** EOARD  
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FPO 09499-0014

**Distribution/Availability Statement:** Approved for public release; distribution is unlimited.

**Abstract:**
This report results from a contract tasking Academy of Sciences as follows: The Grantee will develop new and radically different ideas for a high performance, advanced singlet oxygen generator for driving a supersonic COIL. Various ideas for high performance generator concepts matching up to the above properties should be theoretically considered and the best concept should be developed into the experimental device of advanced SOG. Phase 1: A conceptualisation of the problem including a literature study, theoretical estimation of particular problems, calculations, elaboration of generator concept. At the end of the first year, a consortium of the supervisor, contractor, and Russian and Czech investigators should evaluate the first phase of project solution, and if it is successful, to specify the advanced generator concept that should be further developed.

Phase 2: A design of advanced generator device including detailed technical drawings, purchasing of suitable constructional materials and experimental equipments, construction and testing of individual parts of experimental set-up, construction of small-scale advanced generator.

Phase 3: A testing of performance parameters of the small-scale advanced generator and characterization of its output parameters (O2(1D) yield, residual chlorine and H2O vapor content), and testing this generator with coupling to the laser.

**Subject Terms:** EOARD, Chemical oxygen iodine lasers

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**Security Classification of:**
- a. Report UNCLASS
- b. Abstract UNCLASS
- c. This page UNCLASS

**Limitation of Abstract:** UNCLASS

**Number of Pages:** 56

**Name of Responsible Person:** DONALD J SMITH

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DECLARATION

(1) The Contractor, Institute of Physics of Academy of Sciences of the Czech Republic, hereby declares that, to the best of its knowledge and believes, the technical data delivered herewith under Contract No. FA8655-04-1-3021 is complete, accurate, and complies with all requirements of the contract.

(2) I certify that there were no subject inventions to declare as defined in FAR 52.227-13, during the performance of this Contract.

Date: 31 January 2005

Name and Title of Principal Investigator:          Dr. Jarmila Kodymová

Name and Title of Authorized Official:      Dipl. Ing. Karel Jungwirth, DrSc.
            Director of Institute of Physics AS CR
Objects and goals of developing an advanced SOG

A. Concept investigated in the COIL Laboratory in Samara:

The conceptual designing of the small-scale Rotating Sparger SOG (RSSOG) (30 mmol/s) with next requirements

1. Estimations of hydrodynamic and mass-transfer rates, chlorine utilization and \( \text{O}_2(\Delta) \) yield
   1.1. Basic data and formulas
   1.2. Estimation of RSSOG parameters for achieving \( P_0=100 \) Torr
   1.3. Parameters of RSSOG at \( P_0<100 \) Torr

2. Designing of parameters for the single pass burn down small-scale RSSOG
   \((M(\text{O}_2)=30 \text{ mmole/s}, P_0=100 \text{ Torr})\)

3. Principle constructions of the small-scale RSSOG for production of 30 mmol/s of oxygen flow (close coupling of the generator to nozzle plenum to minimize \( \text{O}_2(\Delta) \) loss and disengagement of gas/liquid)

4. Conclusion

B. Concept(s) investigated in the COIL Laboratory in Prague

Centrifugal Spray Singlet Oxygen Generator (CentSpraySOG)

1. Principle design and operation of CentSpraySOG
2. Estimation of main characteristics of the spray SOG
3. Detailed theoretical description
   3.1. Mass transfer in liquid phase
   3.2. Mass transfer at g/l interface
4. Numerical solution of generator parameters
5. Two-phase nozzles for spray formation
6. Model calculations for chosen nozzles
   6.1. Results of parametric estimations
   6.2. Conclusions from modelling
7. Experimental set-up for measuring a drop size and drop velocity
   7.1. Methods available for these measurements
   7.2. Scheme of device for spray parametric study
8. Design gas/liquid blade separator for CentSpraySOG
   8.1. Geometry of channels between next blades in separator
   8.2. Hydrodynamics of flowing film along blades
   8.3. Settling (disengagement) of droplets
   8.4. Geometry of liquid separator and parameters of liquid film
   8.5. Calculation of pressure loss and singlet oxygen loss in gas channels
9. Design concept of CentSpraySOG device
10. Conclusions of present study on CentSpraySOG

References

Acknowledgment
Objects and goals of developing an advanced SOG

This one-year EOARD grant was awarded to promote the co-operation of COIL Laboratories at Institutes of Physics in Prague, Czech Republic and Samara, Russia on development new concepts of advanced chemical generator of singlet oxygen for a COIL. The generator should have the following characteristics:

1. A high-pressure operation with \(~100\) Torr of \(\text{O}_2\) partial pressure
2. A single BHP pass burn down (probably requiring a dilute BHP to maintain the thermal control and so to limit \(\text{H}_2\text{O}\) vapor)
3. A close coupling of the generator to the nozzle plenum to minimize \(\text{O}_2\left(1\Delta\right)\) losses
4. An efficient disengagement of gas/liquid (conceptually involving the angular acceleration of fluid and gas or both)
5. A scalability to be suitable for the airborne and mobile applications

The investigation within the framework of this grant has been aimed at conceptualisation of the problem including a literature study, theoretical estimation of particular problems, calculations, and designing the concept of the experimental device.
A. The conceptual designing of the small-scale Rotating Sparger SOG (RSSOG) (30 mmole/s) with next requirements

1. Estimations of hydrodynamic and mass-transfer rates chlorine utilization and O$_2$(^1Δ) yield

1.1. Basic data and formulas

The estimation of RSSOG parameters is based on the simplified liquid-gas hydrodynamic and simplified model of the transport phenomena of the bubbler centrifugal apparatus performed in the interim Report 001 of this grant.

The parameters of RSSOG that should be found for achieving the required parameters of the advanced SOG are presented in Table 1.

<table>
<thead>
<tr>
<th>Tab. 1</th>
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<table>
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<th>Tab. 2</th>
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<td>11</td>
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</tbody>
</table>
Tab. 3
Given and calculated intermediate parameters

<table>
<thead>
<tr>
<th></th>
<th>Formula</th>
<th>Description</th>
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<tbody>
<tr>
<td>1</td>
<td>$T=253^\circ K$</td>
<td>Input BHP temperature</td>
</tr>
<tr>
<td>2</td>
<td>$P_0=100$ torr</td>
<td>Total pressure ($O_2 + Cl_2$resid) over BHP layer</td>
</tr>
<tr>
<td>3</td>
<td>$A=0$</td>
<td>Dilution with Helium</td>
</tr>
<tr>
<td>4</td>
<td>$w_r=\omega R$</td>
<td>Peripheral BHP velocity</td>
</tr>
<tr>
<td>5</td>
<td>$G=\omega^2 R$</td>
<td>Centrifugal acceleration</td>
</tr>
<tr>
<td>6</td>
<td>$P=P_0+\omega^2 RH$</td>
<td>Gas pressure at the nozzle outlet (torr)</td>
</tr>
<tr>
<td>7</td>
<td>$\rho_g=(P/RT)(71/(1+A)+A*4/(1+A))$</td>
<td>Gas density at the nozzle outlet ($g/cm^3$)</td>
</tr>
<tr>
<td>8</td>
<td>$N_c=P/RT/(1+A)$</td>
<td>Chlorine concentration at the nozzle outlet (mole/cm$^3$)</td>
</tr>
<tr>
<td>9</td>
<td>$\mu_g=3\times10^{-3}\times A/(1+A)+1.6\times10^{-7}/(1+A)$</td>
<td>Gas viscosity ($g/cm/s$)</td>
</tr>
<tr>
<td>10</td>
<td>$D_c=(1+A)/P(2/300+1/100)^{-1}$</td>
<td>Chlorine diffusion coefficient in $O_2$-He mixture ($cm^2/s$)</td>
</tr>
<tr>
<td>11</td>
<td>$Re=\frac{\rho U_d d_g}{\mu}$</td>
<td>Reynolds number for the gas in nozzle</td>
</tr>
</tbody>
</table>

Tab. 4
Estimation of the hydrodynamic parameters of the rotary bubbler, specific chlorine absorption time and $O_2(\Delta)$ yield parameters

<table>
<thead>
<tr>
<th></th>
<th>Formula</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>$r=r_0\times0.68\left(\frac{\rho_r}{\rho_l}\right)^{0.14}\left(\frac{U_d^2}{d_g}\right)^{0.16}$</td>
<td>Radius of bubble</td>
</tr>
<tr>
<td>2</td>
<td>$U\approx1.43\sqrt{rG}$</td>
<td>Velocity of bubble</td>
</tr>
<tr>
<td>3</td>
<td>$\tau=\frac{1}{C}\sqrt{0.12\times10^{-4}C^2+1.05\times10^{-3}C+0.21-0.35\times10^{-3}}$</td>
<td>Time of bubble formation</td>
</tr>
<tr>
<td>4</td>
<td>$C=\sqrt{G\left(\frac{\rho_g d_g}{d_0}\right)^{0.1}}\frac{Re}{0.06}$</td>
<td>Time of bubble formation</td>
</tr>
<tr>
<td>5</td>
<td>$f=\sqrt{0.25\times C^2+0.71\times C+0.19\times10^6-0.44\times10^3}$</td>
<td>Frequency of bubbles generation</td>
</tr>
<tr>
<td>6</td>
<td>$t_b=\tau+H/U$</td>
<td>Bubble residence time in BHP</td>
</tr>
<tr>
<td>7</td>
<td>$t_{ab}\approx0.6r^2/D_c^2$</td>
<td>Chlorine absorption time due to gas resistance</td>
</tr>
<tr>
<td>8</td>
<td>$t_r=H, r, N_t, D_c$</td>
<td>Chlorine absorption time according to surface reaction</td>
</tr>
<tr>
<td>9</td>
<td>$t_{ab}=\frac{T_{ab}}{N_c/D_c\pi D_h}$</td>
<td>Time of strong $HO_2$ depletion (formation of sheet flame)</td>
</tr>
<tr>
<td>10</td>
<td>$t_{ab}=\frac{T_{ab}}{N_c/D_c\left(\frac{r}{U_d^2}\right)^{03}(2\pi)^{0.5}}$</td>
<td>Time of total chlorine absorption in sheet flame approximation</td>
</tr>
<tr>
<td>11</td>
<td>$U=\frac{h}{\frac{t_{ab}}{U}}\left[\left(\frac{P}{P_0}-1\right)\left(\frac{P}{P_0}-1\right)\right]$</td>
<td>Estimated chlorine utilization for moving bubble (see below)</td>
</tr>
</tbody>
</table>
### Note to Item 11: Estimation of chlorine utilization:

Because of very high initial chlorine pressure (more than 100 torr under BHP layer) and very fast depletion of the \( \text{HO}_2^- \) ions, the chlorine absorption rate will occur according to the flame sheet approximation. In this approximation the chlorine flow rate to bubble surface will be

\[
J_c = \frac{4}{3} N_n D_{\text{th}} \left( \frac{3 U r(x)^3}{2 \pi D_{\text{th}}} \right)^{0.5}
\]

The chlorine absorption from the moving bubble is described by the equation

\[
U \frac{d}{dx} \left( \frac{4 \pi}{3} r(x)^3 N_c \right) = -\pi \frac{4}{3} N_n D_{\text{th}} \left( \frac{3 U r(x)^3}{2 \pi D_{\text{th}}} \right)^{0.5}
\]

Here \( x \) is a distance from the bottom of the BHP layer, \( r(x) \) is the bubble radius.

\[
M_c = f N_n \times 4 \pi r^3 / 3 N_n
\]

Estimated oxygen flux from the bubble layer (mmole/cm²/s)

---

**Table**

<table>
<thead>
<tr>
<th>Item</th>
<th>Equation</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>12</td>
<td>( Y_{\text{der}} \approx 0.9 \exp \left( \frac{-2 N_c}{H N_H - \frac{\pi D_{r} r}{U D_{e} \tau_{e}}} \right) )</td>
<td>Estimation of minimal detachment ( \text{O}_2\left(^1\Delta\right) ) yield</td>
</tr>
<tr>
<td>13</td>
<td>( P_{ct} = \frac{(P+P_0)(A+1)*t_0/4}{Y_{\text{der}}(1+Y_{\text{der}} P_{ct})} )</td>
<td>Average ( P\tau ) parameter due to homogeneous quenching of ( \text{O}_2\left(^1\Delta\right) ) yield</td>
</tr>
<tr>
<td>14</td>
<td>( Y = \frac{Y_{\text{der}}(1+Y_{\text{der}} P_{ct})}{Y_{\text{der}}(1+Y_{\text{der}} P_{ct})} )</td>
<td>Estimation of ( \text{O}_2\left(^1\Delta\right) ) yield</td>
</tr>
<tr>
<td>15</td>
<td>( N_n = (2r)^2 )</td>
<td>Maximum density of chlorine nozzles (1/cm²) to avoid bubble coalescence</td>
</tr>
<tr>
<td>16</td>
<td>( M_c = f N_n \times \frac{4 \pi r^3}{3} N_n )</td>
<td>Estimated oxygen flux from the bubble layer (mmole/cm²/s)</td>
</tr>
</tbody>
</table>
\[
\frac{dn_{\text{c}H}}{dx} = -\frac{N_{\text{H}}}{N_{\text{c}}} \left( \frac{3D_{\text{H}}}{2\pi^3 U} \right)^{0.5} \frac{P_0}{\left( P_0 + \rho G(h - \chi) \right)^{0.5}}
\]

(4)

The solution of this equation gives the estimation of the chlorine utilization

\[
Ut = 2 \frac{N_{\text{H}}}{N_{\text{c}}} \left( \frac{3D_{\text{H}}}{2\pi^3 U} \right)^{0.5} \left( \frac{P}{P_0} - 1 \right) \left( \frac{P}{P - P_0} \right) = \frac{h}{t_\text{sh} U} \left( \frac{P}{P_0} - 1 \right) \left( \frac{P}{P - P_0} \right)
\]

(5)

In this solution the bubble velocity was assumed constant. Indeed \( U = 1.43 (r(x)G)^{0.5} = 1.43 (rG)^{0.5} (P/P(x))^{1/6} \) and weakly depends on \( x \). The formula (5) is valid when calculated \( Ut < 1 \). The case \( Ut >> 1 \) means that the flame sheet approximation is not valid and a surface reaction approximation have to be used. When \( Ut > 1 \), the absorption mode is intermediate between flame sheet approximation and surface reaction approximation. In this case it is necessary to compare values of times \( t_s, t_r \).

**Note to Item 12:** The estimation of the detachment yield in the flame sheet model approximation is

\[
Y_{\text{det}} \approx \frac{\alpha}{\pi D_{\text{c}} D_\alpha} \exp \left( - \frac{2N_{\text{c}}}{H_i N_{\text{H}}} \frac{\pi D_{\text{c}} r}{D_\alpha \tau_\alpha} \right), \quad \frac{\alpha}{\pi D_{\text{c}} D_\alpha} \approx 0.9
\]

where \( t \) is a time of starting of the BHP-\( \text{Cl}_2 \) contact. The most of the oxygen gas is produced near the bubble equator (\( t = r/U \)). Hence the estimation of the \( \text{O}_2(\Delta) \) detachment yield is

\[
Y_{\text{det}} \approx 0.9 \exp \left( - \frac{2N_{\text{c}}}{H_i N_{\text{H}}} \frac{\pi D_{\text{c}} r}{UD_\alpha \tau_\alpha} \right)
\]

**Note to Item 13:** \( \text{Pr} \) is defined as an average for pressure \( (P + P_0) \) and divided by 2 because the \( \text{O}_2(\Delta) \) is monotonically produced during time \( t_r \).

**Note to Item 14:** It is assumed that distance between nozzles should be more than bubble diameter.

1.2. Estimation of RSSOG parameters for achieving \( P_0 = 100 \text{ torr} \).

A one example of the estimations of RSSOG parameters have been performed for the next a priory given input parameters and varied number of the revolutions per second:
Tab. 5

<p>| | | |</p>
<table>
<thead>
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</thead>
<tbody>
<tr>
<td>1</td>
<td>R=3 cm</td>
<td>Radius of RSSOG</td>
</tr>
<tr>
<td>2</td>
<td>d₀=0.02 cm=200 μm</td>
<td>Diameter of holes</td>
</tr>
<tr>
<td>3</td>
<td>H=0.3 cm</td>
<td>Height of BHP layer</td>
</tr>
<tr>
<td>4</td>
<td>w=0</td>
<td>Axial BHP velocity</td>
</tr>
<tr>
<td>5</td>
<td>n-varied</td>
<td>Revolutions per minute (RPM)</td>
</tr>
<tr>
<td>6</td>
<td>U₀=130 m/s</td>
<td>Gas velocity from nozzle</td>
</tr>
<tr>
<td>7</td>
<td>N₁=6 mole/L</td>
<td>BHP molarity (KOH)</td>
</tr>
</tbody>
</table>

The results of calculations according to the formula in Table 4 and for the input parameters in Table 5 are presented in Figs. 1-5. The dependences presented in Figures 1-5 can be explained qualitatively. When the centrifugal acceleration G is not too big the pressure P is close to P₀, and according to eq. (5), the utilization increases with increasing RPS. At higher G, the pressure P is substantially higher than P₀ and the chlorine concentration in the nozzle increases with increasing G. It explains non-monotonic dependence of the utilization, detachment yield and Pt parameter with increasing RPS.

Fig. 1 The initial bubble radius as a function of the rotation rate (RPS)
**Fig. 2** Dependence of the bubble velocity on the number of RPS

**Fig. 3** Dependence of the chlorine utilization on RPS
A lot of calculations have been performed. How the parameters indicated in Figs.1-5 are changed when other parameters in Table 5 are changed:

a) Nozzle diameter. A decrease in the nozzle diameter (lower than 200µm) results in a smaller bubble size, lower bubble velocity and higher utilization. However, simultaneously $P_\tau$ increases and $Y_{det}$ drops. It is necessary to decrease simultaneously the BHP layer height to maintain a high utilization and $O_2(^1\Delta)$ yield.

b) The gas velocity in the nozzles. An increase in the gas velocity results in increasing the bubble size and bubble velocity, which causes the utilization decrease. The maximum density of the nozzles in the bubbler decreases, the frequency of the bubble production decreases, and consequently the maximum oxygen flux doesn’t increase substantially with increasing $U_g$.

c) The height of the BHP layer. With an increase in $H$, the utilization increases. However, increasing $H$ results in the increase in the flight time $t_f$, and the increase in the bottom pressure $P$. This causes the $Y_{det}$ drop and $P_\tau$ increase.

d) The BHP molarity. The decrease in $N_H$ substantially reduces the chlorine utilization and the detachment yield at a high initial chlorine pressure. To attain the required parameters with diluted BHP, it is necessary to decrease the nozzle diameter, gas velocity in the nozzle, the BHP height, and to increase the density of the nozzles.
Several sets of the parameters presented in Table 1 have been used in calculations. We select only those sets of the initial parameters, at which the required output parameters have been achieved. Table 6 presents the estimation of other sets of the input parameters resulting in achieving a high utilization and \( \text{O}_2(1\Delta) \) yield of \(~70\%\) at the output pressure \( P_0=100 \text{ torr} \).

**Tab. 6**

The recommended initial RSSOG parameters for achievement of the high utilization and \( \text{O}_2(1\Delta) \) yield; the radius of the bubbler apparatus is \( R=3 \text{ cm} \)

<table>
<thead>
<tr>
<th>Input parameters</th>
<th>Output parameters</th>
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</thead>
<tbody>
<tr>
<td>( n )</td>
<td>( H )</td>
</tr>
<tr>
<td>( s^{-1} )</td>
<td>( \text{mm} )</td>
</tr>
<tr>
<td>1</td>
<td>60</td>
</tr>
<tr>
<td>2</td>
<td>100</td>
</tr>
<tr>
<td>3</td>
<td>80</td>
</tr>
<tr>
<td>4</td>
<td>100</td>
</tr>
<tr>
<td>5</td>
<td>80</td>
</tr>
</tbody>
</table>

*Note:* For all initial parameters \( t_{fg}<t_{ag} \) and \( t_{dep}<t_{ag} \). Hence the flame sheet model approximation is valid.

The parameters in Table 6 correspond to only very rough estimation of RSSOG parameters. The main parameter for the COIL is the chlorine flux \( M_c \) or oxygen molar flow rate produced from 1 cm² of the bubbler layer. Parameterization showed that a high \( Y_{det}>80\% \) and low \( P_r<0.1 \text{ torr*s} \) could be achieved at \( M_c \leq 1 \text{ mmole/cm}^2/\text{s} \), which is not enough large for the use in a COIL. The main reason is in requirement of the high pressure \( P_0=100 \text{ torr} \). The quenching of \( \text{O}_2(1\Delta) \) and depletion of \( \text{HO}_2^- \) in the

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**Fig. 5** Dependence of the quenching parameter \( P_r \) on the RPS
The bubbler layer is very strong to ensure simultaneously a high utilization and \( Y > 50\% \) at \( M_c > 1 \) mmole/cm\(^2\)/s and \( P_0 = 100 \) torr. Moreover, it is quite difficult to ensure good output parameters at the low BHP molarity. A higher \( M_c \geq 3 \) mmole/cm\(^2\)/s is possible to achieve at lower predicted yield \( Y_{\text{det}} \leq 70\% \) and higher \( P_T > 0.2 \) torr\(\times\)s. So at higher \( M_c \) the predicted \( O_2(1^{\Delta}) \) yield would be lower than 50\%. It seems that the real liquid-gas hydrodynamic and mass transfer phenomena are much more complex. The heating of the BHP results in increasing in the mass transfer coefficients \((D_c, D_H, D_o)\), the decrease in the BHP viscosity. These phenomena have to improve chlorine utilization. The real output parameters of RSSOG can be much better at higher \( M_c \). But only experiment can clear these problems and questions.

1.3. Parameters of RSSOG at \( P_0 < 100 \) torr.

It was found that a decrease in the output pressure \( P_0 \) makes possible to increase the RSSOG specific output flux \( M_c \) and to save a high chlorine utilization and \( O_2(1^{\Delta}) \) yield. Two sets of the input RSSOG parameters at \( P_0 = 20 \) torr are presented in Table 6a.

Tab. 6a
Example of the input-output parameters of the RSSOG at moderate \( P_0 = 20 \) torr

<table>
<thead>
<tr>
<th>Input parameters</th>
<th>Output parameters</th>
</tr>
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<tbody>
<tr>
<td>( n ) s(^{-1})</td>
<td>( H ) mm</td>
</tr>
<tr>
<td>6</td>
<td>80</td>
</tr>
<tr>
<td>7</td>
<td>80</td>
</tr>
</tbody>
</table>

2. Designing of parameters for the single pass burn down small-scale RSSOG \((M(O_2)= 30 \) mmole/s, \( P_0 = 100 \) torr)

Additional specifications:

- \( \eta_w \) – water vapour fraction
- \( P_wi \) – saturated water vapour pressure for the input BHP temperature \( T \)
- \( P_{wsf} \) – saturated water vapour pressure for the output BHP temperature \( T_f \)
- \( Y \) – expected \( O_2(1^{\Delta}) \) yield
- \( Z \) – axial length of the rotary bubbler
- \( N_{Hf} \) – output concentration of the \( HO_2^- \) ions
- \( t_{BHP} \) - BHP residence time in RSSOG
- \( M(O_2) \) - output oxygen molar flow rate
In the single pass burn down RSSOG the output water vapour fraction should be enough low and the BHP consumption high. The estimations made below are based on the results present in the Report 001. We assume that the water fraction $\eta_w=15\%$ to the oxygen concentration does not decrease substantially the COIL gain and power in comparison with $\eta_w=0$. The fraction of the water vapour is given by ((24), Report 001):

$$\eta_w \approx \frac{(1 + A)P_{wsl}(P_{wsl}/P_{wsl}) - 1}{\ln \left( \frac{P_{wsl}}{P_{wsl}} \right)}$$

(6)

The water vapour partial pressure at input BHP temperature is given by

$$P_{wsl}(\text{torr}) = \exp \left( 23.44 - \frac{6051}{T} \right)$$

(7)

The water vapour partial pressure at output BHP temperature is given by

$$P_{wsf}(\text{torr}) = \exp \left( 23.44 - \frac{6051}{T_f} \right)$$

(8)

Assuming $T=253K$, dilution $A=0$, and $P_0=100$ torr, we obtain the required output BHP temperature $T_f \leq 315K$ to maintain $\eta_w \leq 0.15$. The required maximum BHP residence time ($t_{BHP}$) in the RSSOG to ensure $T_f \leq 315K$ and $\eta_w \leq 0.15$ is given by

$$t_{BHP} = \frac{Z}{W} = \frac{T_f - T_i}{M (Q_r + (1 - Y)Q_\Delta)}$$

(9)

At maximum residence time (9) the output $\text{HO}_2^-$ concentration is given by

$$N_{\text{eff}} = N_{\text{eff}} - \frac{2M}{H} t_{BHP}$$

(10)

In (9) and (10), $C_{\text{BHP}}=3.7 \text{ J/cm}^3/\text{K}$ is the volume heat capacity of the BHP, $Q_r = 112 \times 10^3 \text{ J/mole}$ is the heat release in reaction $\text{Cl}_2 + 2\text{HO}_2^- \rightarrow 2\text{Cl}^- + \text{H}_2\text{O} + \text{O}_2(\Delta)$, $Q_\Delta = 94 \times 10^3 \text{ J/mole}$ is the heat release in reaction $\text{O}_2(\Delta) \rightarrow \text{O}_2(\Sigma)$.

For the oxygen molar flow rate $M(\text{O}_2)$ and radius of the bubbler cylinder $R=3$ cm the, length $Z$ of the perforated bubbler cylinder in the axial direction, the BHP axial velocity $w$ and minimal volumetric rate $V_{\text{BHP}}$ are correspondently equal to

$$Z = \frac{M(\text{O}_2)}{M_c 2\pi R}$$

(11)

$$w = \frac{Z}{t_{BHP}} = \frac{M(\text{O}_2)}{M_c 2\pi R t_{BHP}}$$

(12)

$$V_{\text{BHP}} = 2\pi RHw = \frac{M(\text{O}_2)(Q_r + (1 - Y)Q_\Delta)}{C_{\text{BHP}}(T_f - T_i)}$$

(13)
The minimal $V_{\text{BHP}}$ depends only on required oxygen molar flow rate and on $T_f$ (or $P_0$). If $N_{\text{Hf}}$ defined by (10) is very low or less than 0 the BHP volumetric rate should be higher than defined from water vapour management (13). We will suppose in this case that the BHP residence time should be short enough to support $N_{\text{Hf}} \geq \frac{N_{\text{H}}}{2}$ and good chlorine absorption conditions. Hence the maximum BHP residence time is in this case

$$t_{\text{BHP}} = \frac{N_{\text{H}}}{4}(\frac{H}{M_c})$$

Equation (14)

For RSSOG parameters in Tables 6 and 6a, we obtain the required BHP residence time in RSSOG, consumption of BHP alkaline molarity, the axial length, axial velocity and volumetric rate of the BHP. Hence the final designing parameters and expected output parameters are presented in Table 7.

### Tab. 7

The preliminary designing parameters and expected output parameters for $M(O_2)=30 \text{ mmole/s}$ (explanation in Fig.6)

<table>
<thead>
<tr>
<th>Item</th>
<th>$n$ s$^{-1}$</th>
<th>$H$ mm</th>
<th>$d_0$ µm</th>
<th>$U_g$ m/s</th>
<th>$N_H$ mole/L</th>
<th>$M_c$ mmole/cm$^2$/s</th>
<th>$t_{\text{BHP}},$s</th>
<th>$Z$ cm</th>
<th>$w$ cm/s</th>
<th>$V_{\text{BHP}}$ L/s</th>
<th>$N_{\text{Hf}}$ mole/L</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>60</td>
<td>3</td>
<td>200</td>
<td>130</td>
<td>6</td>
<td>$\sim$1</td>
<td>0.5</td>
<td>1.6</td>
<td>3.2</td>
<td>0.018</td>
<td>2.7</td>
</tr>
<tr>
<td>2</td>
<td>100</td>
<td>1.5</td>
<td>100</td>
<td>130</td>
<td>6</td>
<td>$\sim$1</td>
<td>0.25</td>
<td>1.6</td>
<td>6.4</td>
<td>0.018</td>
<td>2.7</td>
</tr>
</tbody>
</table>

In Items 3-4 the BHP volumetric flow rate should be higher than defined from (13) or $t_{\text{BHP}}$ should be defined from (14) to keep $N_{\text{Hf}} \geq \frac{N_{\text{H}}}{2}$ mole/litre

| 3    | 80           | 6      | 300      | 130      | 6            | $\sim$3            | 0.3              | 0.53 | 1.8    | 0.02            | 3              |
| 4    | 80           | 1      | 50       | 100      | 3            | $\sim$0.5          | 0.15             | 3.2  | 21.3   | 0.04            | 1.5             |
| 5    | 100          | 1      | 50       | 50       | 2            | $\sim$0.5          | 0.1             | 3.2  | 32     | 0.06            | 1              |

Input parameters for the moderate pressure RSSOG $(T=253K, P_0=20 \text{ torr}, \text{Table 6a})$

<table>
<thead>
<tr>
<th>Item</th>
<th>$n$ s$^{-1}$</th>
<th>$H$ mm</th>
<th>$d_0$ µm</th>
<th>$U_g$ m/s</th>
<th>$N_H$ mole/L</th>
<th>$M_c$ mmole/cm$^2$/s</th>
<th>$t_{\text{BHP}},$s</th>
<th>$Z$ cm</th>
<th>$w$ cm/s</th>
<th>$V_{\text{BHP}}$ L/s</th>
<th>$N_{\text{Hf}}$ mole/L</th>
</tr>
</thead>
<tbody>
<tr>
<td>6</td>
<td>80</td>
<td>5</td>
<td>500</td>
<td>200</td>
<td>6</td>
<td>$\sim$3</td>
<td>0.14</td>
<td>0.53</td>
<td>3.8</td>
<td>0.036</td>
<td>4.3</td>
</tr>
<tr>
<td>7</td>
<td>80</td>
<td>4</td>
<td>300</td>
<td>100</td>
<td>3</td>
<td>$\sim$2</td>
<td>0.1</td>
<td>0.8</td>
<td>8</td>
<td>0.06</td>
<td>2</td>
</tr>
</tbody>
</table>

The efficiency of the BHP consumption at high $P_0$ is close to 50%. At moderate pressure $P_0=20 \text{ torr}$ the consumption of the BHP is lower. The most suitable input parameters are in Item 3 at $P_0=100 \text{ torr}$ and Item 6 for $P_0=20 \text{ torr}$.

Fig. 6 Designing parameters of RSSOG
3. Principle constructions of the small-scale RSSOG for production of 30 mmole/s of oxygen flow (close coupling of the generator to nozzle plenum to minimize O₂(1∆) loss and disengagement of gas/liquid)

The volumetric rate of the gas over BHP bubbler layer is

\[ V_{\text{gas}} = \frac{M(O2)RT}{P_0} \]  

(15)

\( R_g = 8.31 \text{ J/mole/K} \) is the universal gas constant. The gas transport time from the BHP bubbler layer to the throat of the nozzle bank is equal to

\[ t_{\text{trans}} = \frac{V_0}{V_{\text{gas}}} = \frac{V_0P_0}{M(O2)R_gT}, \]  

(16)

\( V_0 \) is the volume of the transport duct between the BHP surface and the nozzle throat.

The so-called quenching parameter \( \rho \) due to reactions \( O_2(1\Delta) + O_2(1\Delta) \rightarrow O_2(1\Sigma) + O_2(3\Sigma) \) is equal to

\[ \rho t_{\text{trans}} = \frac{P_0V_0}{V_{\text{gas}}} = \frac{V_0P_0^2}{M(O2)R_gT}, \]  

(17)

Let’s require \( (P_0t_{\text{trans}}) < \chi_{\text{max}} = 0.1 \text{ torr}\times\text{s} \). Hence the maximum volume of the transport duct

\[ V_0 < \frac{2}{P_0} \frac{M(O2)}{P_0} \]  

(18)

For the input parameters presented in Table 7, the maximum volumes of the transport duct between the bubbler BHP layer and the nozzle throat are presented in Table 8.

Tab. 8

(Items 1-7 are the same like in Tables 6 and 6a)

<table>
<thead>
<tr>
<th></th>
<th>Output pressure P₀, torr</th>
<th>Axial length of RSSOG Z, cm</th>
<th>Chlorine flux through bubbler surface Mᵣ, mmole/cm²/s</th>
<th>Required maximum volume of transport duct from BHP surface to nozzle throat V₀, cm³</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>100</td>
<td>1.6</td>
<td>1</td>
<td>4.8</td>
</tr>
<tr>
<td>2</td>
<td>100</td>
<td>1.6</td>
<td>1</td>
<td>4.8</td>
</tr>
<tr>
<td>3</td>
<td>100</td>
<td>0.53</td>
<td>3</td>
<td>4.8</td>
</tr>
<tr>
<td>4</td>
<td>100</td>
<td>3.2</td>
<td>0.5</td>
<td>4.8</td>
</tr>
<tr>
<td>5</td>
<td>100</td>
<td>3.2</td>
<td>0.5</td>
<td>4.8</td>
</tr>
<tr>
<td>6</td>
<td>20</td>
<td>0.53</td>
<td>3</td>
<td>120</td>
</tr>
<tr>
<td>7</td>
<td>20</td>
<td>0.8</td>
<td>2</td>
<td>120</td>
</tr>
</tbody>
</table>

Table 8 shows that the required transport duct volume is too small for Items 1-5. It is quite difficult to imagine an assembling of RSSOG with \( R=3 \text{ cm} \) and \( Z\approx1 \text{ cm} \), and with the nozzle bank keeping the plenum pressure \( P_0=100 \text{ torr} \) and keeping \( O_2(1\Delta) \) yield (Items 1-5).
Much more promising is the scheme of COIL with output oxygen pressure $P_0=20$ torr (Items 6-7). Also the COIL schemes with RSSOG parameters according to Items (1-5) with output pressure $P_0=100$ torr can be realized using oxygen pressure discharge from the output pressure $P_0=100$ torr to the plenum pressure $P_{\text{plen}}=20$ torr. The discharge of the pressure can be realized immediately after oxygen outlet from BHP. In this case the required maximum transport volume is equal to 120 cm$^3$ (like for items 6-7) and large enough for the arrangement of the nozzle bank.

Scheme 1

The principle scheme of RSSOG with output and plenum oxygen pressure 20 torr is presented in Fig.7. The RSSOG can be scaled in Z and R directions. The appropriate changes of parameters $w$, $V_{\text{BHP}}$ should be made according to formulas (11)-(14). For example, increasing $Z$ twice forces an increase in the axial velocity $w$ and the BHP volumetric rate twice. The principle assembling of three or several RSSOG is presented in Fig.8.

Fig. 7 Principle scheme of RSSOG
Let’s present some technical characteristics of the assembling of several RSSOG. The oxygen plenum flux (in front of the nozzle bank) is \( M_{O_2} = n M_c 2\pi R / L \), where \( n \) is the number of RSSOG, \( L \) is the length of assembling on \( n \) RSSOG. For \( M_c = 3 \) (mmole/s)/cm\(^2\) the oxygen plenum flux of the order of 7 (mmole/s)/cm\(^2\) can be generated by this set of small scale RSSOG. The ratio of the total oxygen molar flow rate to the BHP volumetric rate of the order of 1.5 (mole/s)/(litre/s) for Item 3 and 0.8 (mole/s)/(litre/s) for Item 6 can be achieved.
Scheme 2

The principle scheme with output pressure 100 torr and plenum pressure 20 torr is presented in Fig. 9 and is similar to the scheme in Fig. 8 with the difference that a perforated wall is installed over the BHP surface to discharge the oxygen pressure from 100 torr to 20 torr.

Fig. 9  Principle scheme of RSSOG with oxygen pressure discharge
Scheme 3

The assembling of several RSSOG in vertical Z direction is presented in Fig. 10. In this scheme, the length Z is of the order of several cm and axial velocity w of the order of several ten centimetres per second. The worked-out BHP drains to the common collector.

Fig. 10 Principle scheme of RSSOG with vertical scaling
4. Conclusions

The conception of the centrifugal bubble SOG is based on understanding that the mass transfer phenomena for these conditions are much faster than in any another type mass-transfer apparatus. Hence very efficient production of the oxygen flow with high \( \text{O}_2(\Delta) \) yield and chlorine utilization is possible. The estimation of the gas-liquid hydrodynamic of centrifugal bubble SOG is based on the literature data reviewed in Report 001 of this grant. The estimation of chlorine utilization and \( \text{O}_2(\Delta) \) production is based on the modern understanding of the mass-transfer phenomena in the chlorine-BHP system. Nevertheless, we understand that all estimations are very rough because the real hydrodynamics and the mass-transfer processes in the centrifugal bubbler SOG are much more complex. Very simplified model of the processes described above gives a very approximate parameters of RSSOG to achieve required output parameters and suitable for COIL application. According to these estimations, the oxygen flux \( M_{\text{O}_2} \sim 3 \) (mmole/s) per 1 cm\(^2\) of the bubbler surface can be produced at the output oxygen pressures 20÷100 torr. A possible production of higher oxygen flux (with high \( \text{O}_2(\Delta) \) yield, high utilization and pressure >20 torr) is a question. For \( M_{\text{O}_2} < 3 \) mmole/s and high \( P_0 \), the transport volume duct between RSSOG and the nozzle ban (NB) should be very small to save \( \text{O}_2(\Delta) \), so it is quite difficult to invent efficient RSSOG + nozzle bank system. The discharge of oxygen plenum pressure to 20 torr would make possible to solve the problem of oxygen transport from the RSSOG to the NB. Using of diluted BHP with low KOH (NaOH, LiOH) concentration substantially decreases the efficiency of chlorine utilization and very hardly designed parameters of RSSOR (a small nozzle diameter, a high centrifugal acceleration) are required. Several schemes of RSSOG and scaling procedures are proposed in this Report. The RSSOG can work in the single pass burn down with efficient BHP consumption. The predicted ratio of chlorine molar flow rate to BHP volumetric rate is of the order of 1(mole/s)/(litre/s) in the single pass burn down mode. This ratio is much higher than for another SOG types. The scaling of RSSOG is evident and clear. A plenum oxygen flux in front of the large scale NB of the order of 7 mmole/cm\(^2\)/s can be produced by assembling several RSSOG.
B. Concept(s) investigated in the COIL Laboratory in Prague

Two concepts of the advanced generator of singlet oxygen were proposed employing a centrifugal force for gas/liquid disengagement: the Centrifugal Film SOG (CentFilmSOG) and Centrifugal Spray SOG (CentSpraySOG). They were described in the interim Report 001 submitted to the EOARD on 31 July 2004. Theoretical estimations of both concepts showed that the CentSpraySOG would better fulfill requirements given in Objects and Goals of this investigation, and therefore, our recent research has been focused on more detailed computational modeling of parameters of this generator, and designing devices for its experimental investigation.

**Centrifugal Spray Singlet Oxygen Generator (CentSpraySOG)**

This concept is based on principle of the efficient generation of singlet oxygen in mixture of chlorine and BHP spray (mist), from which liquid droplets are efficiently separated by a centrifugal force within very short time. The mist of small-diameter droplets of tens microns provides a large specific surface area and a short diffusion path of HO2⁻ ions from the bulk BHP, resulting in substantially better BHP utilization than in any other types of chemical generators (jet-type, disk-type). Calculations performed at the USAF by Blauer, et al. [1] on the aerosol generator showed that the generator operating at moderate O₂(¹Δ) pressures (~20 Torr) and with a high O₂(¹Δ) yield (~70%) may be designed with very restrictive dimensions to limit a time of the two-phase flow below a few milliseconds. Our new proposed concept of CentSpraySOG employing a combination of the spray SOG and centrifugal separator of both phases may overcome this problem.

1. Principle design and operation of CentSpraySOG

The proposed concept is schematically shown in Fig. 1. Operation of the device can be simply described by the following processes:

- An atomizing nozzle (Fig. 2) produces a spray (mist) of BHP droplets by means of pressurized Cl₂/He (N₂) gas mixture.
- Droplets of partially exhausted BHP are separated from g/l mixture by a centrifugal force of the rotating blade separator within a few milliseconds (Fig. 3).
• A liquid-free gas (containing $\text{O}_2(^1\Delta)$, residual $\text{Cl}_2$, buffer gas and residual water vapor) leaves the generator through the exit coupled to the laser nozzle throat, while the liquid is collected in the bottom tank.

![Fig. 1 Scheme of principle of CentSpraySOG device](image)

**Fig. 1** Scheme of principle of CentSpraySOG device

![Fig. 2 Scheme of two-phase spray nozzle](image)

![Fig. 3 Scheme of centrifugal separator](image)

**Fig. 2** Scheme of two-phase spray nozzle  
**Fig. 3** Scheme of centrifugal separator

### 2. Estimation of main characteristics of the spray SOG

Parametric estimation of the spray generator was performed for conditions listed in **Table 1**.
Tab. 1
Main parameters of the reaction system

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Labeling</th>
<th>Magnitude</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gas entering the generator</td>
<td>Cl₂ + He</td>
<td>1 : 4</td>
</tr>
<tr>
<td>Total generator pressure</td>
<td>P_tot</td>
<td>40 kPa (300 torr)</td>
</tr>
<tr>
<td>Gas temperature</td>
<td>T</td>
<td>260 K</td>
</tr>
<tr>
<td>BHP droplet radius</td>
<td>a</td>
<td>10 µm</td>
</tr>
<tr>
<td>Diffusion coefficient of chlorine in gas</td>
<td>D(CG)</td>
<td>3.3 x 10⁻⁵ m² s⁻¹</td>
</tr>
<tr>
<td>Diffusion coefficient of chlorine in BHP</td>
<td>D(CL)</td>
<td>5 x 10⁻⁶ m² s⁻¹</td>
</tr>
<tr>
<td>Diffusion coefficient of HO₂⁻ in BHP</td>
<td>D(HL)</td>
<td>5 x 10⁻⁶ m² s⁻¹</td>
</tr>
<tr>
<td>Diffusion coefficient of oxygen in BHP</td>
<td>D(AL)</td>
<td>5 x 10⁻⁶ m² s⁻¹</td>
</tr>
<tr>
<td>Initial HO₂⁻ concentration in BHP</td>
<td>c(HL₀)</td>
<td>5 kmol m⁻³</td>
</tr>
<tr>
<td>Initial Cl₂ concentration in gas</td>
<td>c(CG₀)</td>
<td>3.7x10⁻³ kmol m⁻³</td>
</tr>
<tr>
<td>Rate constant of reaction Cl₂ + HO₂⁻</td>
<td>k_p</td>
<td>5x10⁸ m³ kmol⁻¹ s⁻¹</td>
</tr>
<tr>
<td>Striking Cl₂ probability at BHP surface</td>
<td>γ_C</td>
<td>0.0075</td>
</tr>
<tr>
<td>Striking O₂(¹Δg) probability at BHP surface</td>
<td>γ_A</td>
<td>0.003</td>
</tr>
<tr>
<td>Henry constant of chlorine</td>
<td>H_C</td>
<td>1</td>
</tr>
<tr>
<td>O₂(¹Δg) lifetime in BHP</td>
<td>τ_A</td>
<td>2 x 10⁻⁵ s</td>
</tr>
<tr>
<td>Rate constant of O₂(¹Δg) dimolar reactions in (g)</td>
<td>k_QG</td>
<td>3.7x10⁴ m³ kmol⁻¹ s⁻¹</td>
</tr>
</tbody>
</table>

The calculations of characteristic times of partial steps in the whole process, i.e., the characteristic time of HO₂⁻ depletion in droplet controlled by interfacial crossing or reactive diffusion in liquid, the characteristic time of Cl₂ depletion, the estimation of O₂(¹Δg) detachment yield and O₂(¹Δg) loss in the gas phase, were presented in detail in the interim report of this grant. Below are summarized therefore only conclusions from these estimations.

- The process of O₂(¹Δg) generation in the proposed conditions is controlled first by chlorine absorption (interfacial crossing) (t < 0.3 ms).
- After this first period, the process is controlled by HO₂⁻ diffusion from the droplet interior.
- Chlorine is nearly exhausted within 3 ms.
- ~ 40% of HO₂⁻ ions of BHP is consumed within this time.
- The O₂(¹Δg) yield at the generator exit decreases with time from 0.73 (after 1 ms) to 0.50 (after 4 ms).
3. Detailed theoretical description of CentSpraySOG

Tab. 2
List of main symbols

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>a</td>
<td>Radius of droplet</td>
</tr>
<tr>
<td>b</td>
<td>Radius of gas volume surrounding the droplet</td>
</tr>
<tr>
<td>( c_{\text{g}} ) (i≡C,H,A,f−G,L)</td>
<td>Molar concentration of HO(_2^\cdot), Cl(_2), and O(_2) in gas and liquid phase</td>
</tr>
<tr>
<td>( c_{\text{Hlg}}, c_{\text{CG0}} )</td>
<td>Initial (bulk) concentration of HO(_2^\cdot) and Cl(_2)</td>
</tr>
<tr>
<td>( \bar{c}<em>{\text{CG}}, \bar{c}</em>{\text{MG}} )</td>
<td>Averaged concentration of Cl(_2) and O(_2) in the gas phase</td>
</tr>
<tr>
<td>( D_{ij} ) (i≡C,H,A,f−G,L)</td>
<td>Diffusion coefficient of species in gas or liquid phase</td>
</tr>
<tr>
<td>t</td>
<td>Time since droplets escaped from the nozzle</td>
</tr>
<tr>
<td>( H_i = \left( \frac{c_{\text{g}}}{c_{\text{lg}}} \right)_{\text{eq}} )</td>
<td>Dimensionless Henry constant of species i</td>
</tr>
<tr>
<td>( \alpha_i = 1 / 4 v_{t,i} \gamma_i )</td>
<td>Mass transfer coefficient of species i across g/l interface</td>
</tr>
<tr>
<td>( \gamma_i )</td>
<td>Stricking probability of species i</td>
</tr>
<tr>
<td>( v_{t,i} )</td>
<td>Thermal velocity of species i</td>
</tr>
<tr>
<td>r</td>
<td>Radial coordinate from the center of droplet</td>
</tr>
<tr>
<td>U_i</td>
<td>Utilization</td>
</tr>
</tbody>
</table>

A volumetric fraction of the liquid in the aerosol can be expressed as

\[
\lambda = \left( 1 + \frac{u c_{\text{Hlg}}}{2 c_{\text{CG0}}} \right)^{-1},
\]

where \( u \equiv U_i / U_c \). If the process runs for an infinitely long time the ultimate concentrations for the case of initial stoichiometry, i.e.

\[
2 V_G c_{\text{CG0}} = V_L c_{\text{Hlg}}
\]

for \( u < 1 \) would be

\[
c_{\text{Hlg}} = (1 - u)c_{\text{Hlg}}, \quad c_{\text{CG0}} = 0,
\]

and for \( u > 1 \):

\[
c_{\text{Hlg}} = 0, \quad c_{\text{CG0}} = (1 - 1/u)c_{\text{CG0}}.
\]

3.1. Mass transfer in liquid phase

A temporal profile of the HO\(_2^\cdot\) concentration in the liquid can be considered for two limiting cases.

3.1.1. Initial (film) stage

In the initial stage, the diffusion-reaction kinetics is limited by molecular transport at the interface. It can be assumed that the flux of HO\(_2^\cdot\) towards the phase interface is constant in time. The corresponding solution is then (adaptation of eq. 1 in [2], p. 242)
where \( \alpha_n, n = 1, 2, \ldots \), are the positive roots of

\[
\tan \alpha = \alpha, \quad \text{or} \quad \alpha \cos \alpha - \sin \alpha = 0,
\]

and \( j_H \) is the molar flux of HO\(_2\). The first root of eq. (6) is \( \alpha = 4.4934 \). Higher roots were found using the Newton-Raphson method with \( \alpha_n(0) = \alpha_{n+1} + \pi \). The iteration of more general form of this equation will be discussed later.

Using dimensionless variables \( y \equiv r / a \) and \( \tau \equiv t D_H / a^2 \), eq. (5) can be written as

\[
c_{HHL} = c_{HHL,0} \left( 1 - \phi_H \left[ 3\tau + \frac{5y^2}{10} - 2 \sum_{n=1}^{\infty} \frac{\sin(\alpha_n y)}{\alpha_n^2 \sin \alpha_n} \exp(-\alpha_n^2 \tau) \right] \right),
\]

where

\[
\phi_H = \frac{j_H a}{D_H c_{HHL,0}}.
\]

The surface concentration \( \gamma = 1 \) is

\[
c_{HHL} = c_{HHL,0} \left( 1 - \phi_H \left[ 3\tau + \frac{1}{5} - 2 \sum_{n=1}^{\infty} \frac{1}{\alpha_n^2} \exp(-\alpha_n^2 \tau) \right] \right) = c_{HHL,0} \{1 - \phi_H \vartheta_H(\tau)\}
\]

The significance of the sum of exponentials decreases sharply with growing \( \tau \). For \( \tau = 0.1 \), it contributes less than 3%.

The mean concentration averaged from \( r = 0 \) to \( r = a \) is

\[
\bar{c}_{HHL} = \frac{2}{\pi a^3} \int_0^a c_{HHL} 4\pi r^2 dr = c_{HHL,0} \left[ 3j_H t \right] a.
\]

If \( \sqrt{D_H t} \ll a \), the influence of curvature can be neglected and the case can be treated as diffusion into a semi-infinite solid (adapt. eq. 7 in [2], p. 75):

\[
c_{HHL} = c_{HHL,0} - 2j_H D_H \left[ \frac{D_H t}{\pi} \exp \left( -\frac{x^2}{4D_H t} \right) - \frac{x}{2} \text{erfc} \left( \frac{x}{2\sqrt{D_H t}} \right) \right],
\]

where \( x = a - r \) and

\[
\text{erfc} \equiv \frac{2}{\sqrt{\pi}} \int_0^x e^{-\xi^2} d\xi.
\]

### 3.1.2. Final (depleted surface) stage

In the final stage, the diffusion-reaction kinetics is limited by the transport of HO\(_2\) towards the phase interface. We assume \( c_{HHL} \ll c_{HHL,0} \), which allows an approximation \( c_{HHL,0} \approx 0 \). An adaptation of eq. 4 in [2], p. 233 gives
\[ c_{H,t} = -c_{H,0} \sum_{n=1}^{\infty} \frac{2a}{n\pi} \left\{ (-1)^n \sin \left( \frac{n\pi r}{a} \right) \exp \left( -\frac{D_a n^2 \pi^2 t}{a^2} \right) \right\}. \]  

(13)

The average concentration at time \( t \) is given as (adapt. of eq. 8. in [2], p. 234)

\[ \overline{c}_{H,t} = c_{H,0} \sum_{n=1}^{\infty} \frac{1}{n^2} \exp \left( -\frac{D_a n^2 \pi^2 t}{a^2} \right). \]  

(14)

The flux at the surface (\( r = a \)) is given as

\[ j_H = -D_a \frac{\partial \overline{c}_{H,t}}{\partial x} = \frac{D_a c_{H,0}}{a} \sum_{n=1}^{\infty} \exp \left( -\frac{D_a n^2 \pi^2 t}{a^2} \right). \]  

(15)

3.1.3. Matching the initial and final solution

An approximation for the diffusion kinetics in the whole domain can be obtained by combining the initial and final solutions. The initial and final solutions are matching at the time \( t_d \), for which two conditions are satisfied: (i) the fluxes are equal, and (ii) the mean concentrations are equal. In order to achieve that solution, a start of the final solution is postponed by a time interval \( \Delta t \), \((\Delta t < t_d)\). Parameters \( t_d \) and \( \Delta t \) are adjusted to satisfy the above two conditions.

The computation is done as follows. We assume that the initial flux is given, and then the dimensionless form can be used

\[ \phi_{H,ini} = \frac{j_{H,ini}}{D_a c_{H,0}}. \]  

(16)

The mean concentration from the initial solution at the matching point is

\[ \overline{c}_{H,ini,d} = c_{H,0} \left( 1 - \frac{3D_a t_d}{a^2} \phi_{H,ini} \right), \]  

(17)

and the dimensionless flux computed from the final solution is given as

\[ \phi_{H,fin,d} = \frac{j_{H,fin,d}}{D_a c_{H,0}} = 2 \sum_{n=1}^{\infty} \exp \left[ -n^2 \pi^2 \frac{D_a}{a^2} (t_d - \Delta t) \right]. \]  

(18)

By setting \( \phi_{H,fin,d} = \phi_{H,ini,d} \), eq. (17) can be iterated to obtain \((t_d - \Delta t)\). The mean concentration at the matching point, computed from the initial solution, is given as

\[ \overline{c}_{H,ini,d} = c_{H,0} \left[ \frac{6}{\pi} \sum_{n=1}^{\infty} \frac{1}{n} \exp \left[ -n^2 \pi^2 \frac{D_a}{a^2} (t_d - \Delta t) \right] \right] = c_{H,0} \phi_{H,fin}(t_d - \Delta t). \]  

(19)

The right-hand side can be evaluated because \((t_d - \Delta t)\) is already known.

By setting \( \overline{c}_{H,fin,d} = \overline{c}_{H,ini,d} \), we then have

\[ \overline{c}_{H,fin} \approx \overline{c}_{H,ini} \]  

(20)
\[ t_d = a \cdot \frac{1 - \theta_{in}(t_d - \Delta t)}{3D_H \Phi_{H,\text{in}}} \quad . \tag{20} \]

The delay time is then obtained as \( \Delta t = t_d - (t_d - \Delta t) \).

### 3.2. Mass transport at g/l interface

#### 3.2.1. Transport across the interface

It is assumed that temperature at both sides of the phase interface is the same. The net flux of the species \( i \) from the gas phase to the liquid phase is

\[ j_{\text{is}} = \alpha_i (c_{\text{isG}} - c_{\text{isL}}^G) = \frac{1}{4 \gamma_i v_{ij}} (c_{\text{isG}} - H c_{\text{isL}}) \quad . \tag{21} \]

#### 3.2.2. Diffusion-reaction kinetics of Cl\(_2\)

We use here the coordinate \( x = r - a \). The diffusion-reaction equation is

\[ \frac{\partial c_{\text{cl}}}{\partial t} = \frac{D_{\text{cl}}}{\partial x^2} - k_r c_{\text{isL}} c_{\text{cl}} \approx 0 \quad . \tag{22} \]

Solution of this equation can be found when approximating \( c_{\text{isL}}(x) \approx c_{\text{isL}} \) (i.e. the concentration variation of \( \text{HO}_2^- \) is much smaller then of Cl\(_2\))

\[ c_{\text{cl}} = c_{\text{clL}} \exp \frac{x}{X_p} \quad , \tag{23} \]

where

\[ X_p = \sqrt{\frac{D_{\text{cl}}}{k_r c_{\text{isL}}} \quad . \tag{24} \]

The flux of Cl\(_2\) is given by

\[ j_{\text{cl}} = -D_{\text{cl}} \frac{\partial c_{\text{cl}}}{\partial x} = -D_{\text{cl}} \frac{c_{\text{clL}}}{X_p} \exp \frac{x}{X_p} = -c_{\text{clL}} \sqrt{D_{\text{cl}} k_r c_{\text{isL}}} \exp \frac{x}{X_p} \quad , \tag{25} \]

and at the surface

\[ j_{\text{clS}} = -D_{\text{cl}} \frac{c_{\text{clL}}}{X_p} = -c_{\text{clL}} \sqrt{D_{\text{cl}} k_r c_{\text{isL}}} \quad . \tag{26} \]

Combining the equations (26) and (21), we obtain

\[ j_{\text{clS}} = -\frac{\alpha_i c_{\text{isG}}}{1 + \alpha_i H c_{\text{isL}} / D_{\text{cl}}} \quad . \tag{27} \]

#### 3.2.3. Diffusion-reaction kinetics of O\(_2\) (\( ^1\Delta \))

The second Fick’s law for singlet oxygen has the form

\[ \frac{\partial c_{\text{sl}}}{\partial t} = D_{\text{sl}} \frac{\partial^2 c_{\text{sl}}}{\partial x^2} + k_r c_{\text{isL}} c_{\text{cl}} - k_{\text{gs}} c_{\text{sl}} \quad . \tag{28} \]

In the steady state, the left side of this equation is nearly zero.
Using (23), setting $c_{\text{HL}}(x) \approx c_{\text{HL}}$, and the condition of $c_{\text{HL}}(x \to \infty) = 0$, we can finally obtain by integration (28)

$$c_{\text{ML}} = c_{\text{ML}}X \exp(x / X_p) + (c_{\text{ML}} - c_{\text{ML}}X) \exp(x / X_{QL}),$$  \hspace{1cm} (29)

where

$$X_{QL} = \sqrt{D_{\text{ML}} / k_{\text{QL}}},$$  \hspace{1cm} (30)

$$X = \frac{D_{\text{CL}}}{D_{\text{ML}}} \left(\left(X_p / X_{QL}\right)^2 - 1\right)^{-1}.$$  \hspace{1cm} (31)

The corresponding flux of O$_2$(1)$\Delta$ is

$$j_{\text{ML}} = -D_{\text{ML}} \frac{\partial c_{\text{HL}}}{\partial x} = -D_{\text{ML}} \left(\frac{c_{\text{CL}}X}{X_p} \exp\left(\frac{x}{X_p}\right) + \frac{c_{\text{ML}} - c_{\text{ML}}X}{X_{QL}} \exp\left(\frac{x}{X_{QL}}\right)\right).$$  \hspace{1cm} (32)

Especially for the phase interface $(x = 0)$, the flux is then

$$j_{\text{ML}} = D_{\text{CL}} c_{\text{CL}} \frac{1}{X_p} \frac{1}{1 + X_p / X_{QL}} - D_{\text{ML}} c_{\text{ML}} \frac{1}{X_{QL}}.$$  \hspace{1cm} (33)

and

$$j_{\text{ML}} = -\alpha_{\text{L}} (c_{\text{ML}} - H_{\text{L}} c_{\text{ML}}).$$  \hspace{1cm} (34)

By combining eqs. (33) and (34), we obtain

$$j_{\text{ML}} = \left(\frac{D_{\text{CL}} c_{\text{CL}}}{X_p} \frac{1}{1 + X_p / X_{QL}} - D_{\text{ML}} c_{\text{ML}} \frac{1}{X_{QL}}\right) \left(1 + \frac{X_\Delta}{X_{QL}}\right).$$  \hspace{1cm} (35)

where

$$X_\Delta = \frac{D_{\text{ML}}}{\alpha_{\text{L}} H_{\text{L}}}.$$  \hspace{1cm} (36)

The length $X_\Delta$ can be interpreted as a thickness of hypothetical additional layer of liquid, through which O$_2$(1)$\Delta$ must diffuse.

Using eq. (35)

$$j_{\text{ML}} = \left(\frac{-j_{\text{CL}}}{1 + X_p / X_{QL}} - D_{\text{ML}} c_{\text{ML}} \frac{1}{X_{QL}} H_{\text{L}}\right) \left(1 + \frac{X_\Delta}{X_{QL}}\right).$$  \hspace{1cm} (37)

3.2.4. Simplified evolution equations for Cl$_2$ and O$_2$(1)$\Delta$ concentrations in gas

Because the diffusion of Cl$_2$ and O$_2$(1)$\Delta$ in the gas phase is fast in comparison with the HO$_2^-$ diffusion in the liquid, it is possible to ignore the spatial gradient of their concentrations:

$$\bar{c}_{\text{CG}} \approx c_{\text{CG}},$$

$$\bar{c}_{\text{MG}} \approx c_{\text{MG}}.$$  \hspace{1cm} (38)

Then the evolution of chlorine concentration is given as
\[
\frac{4}{3} \pi (b^3 - a^3) \frac{d\bar{c}_{CG}}{dt} = 4\pi a^2 j_{cs}, \tag{39}
\]
\[
\frac{d\bar{c}_{CG}}{dt} = \frac{1}{a} \frac{3\lambda}{1 - \lambda} j_{cs}. \tag{40}
\]

It follows from the stoichiometry that
\[
\tau_{CG}(t) = c_{CG0} + \frac{\dot{V}_i}{2\dot{V}_G} [\tau_{HL}(t) - c_{HL0}] = c_{CG0} + \frac{\lambda}{2(1 - \lambda)} [\tau_{HL}(t) - c_{HL0}]. \tag{41}
\]

The evolution of \(O_2(\Delta)\) concentration is given as
\[
\frac{4}{3} \pi (b^3 - a^3) \frac{d\bar{c}_{SG}}{dt} = 4\pi a^2 j_{\Delta} = \frac{4}{3} \pi (b^3 - a^3) k_{GG}\bar{c}_{SG}^2, \tag{42}
\]
\[
\frac{d\bar{c}_{SG}}{dt} = \frac{1}{a} \frac{3\lambda}{1 - \lambda} j_{\Delta} - k_{GG}\bar{c}_{SG}^2. \tag{43}
\]

For constant flux \(j_{\Delta}\) and initial condition \(c_{SG}(t = 0) = 0\) the equation (43) can be integrated:
\[
\bar{c}_{SG}(t) = \frac{1}{k_{GG}t_{\lambda}} \tanh \frac{t}{t_{\lambda}}, \tag{44}
\]
where the characteristic time is
\[
t_{\lambda} = \sqrt{\frac{(1 - \lambda)a}{3j_{\Delta}k_{GG}}}. \tag{45}
\]

Using eq. (43) and (37), we obtain
\[
\frac{d\bar{c}_{SG}}{dt} = \frac{1}{a} \frac{3\lambda}{1 - \lambda} \left( -\frac{j_{GL}}{1 + X_p/X_{GL}} - \frac{D_{SL}c_{SL}}{X_{GL}H_{\lambda}} \right) - k_{GG}\bar{c}_{SG}^2. \tag{46}
\]

The last equation can be written as
\[
\frac{d\bar{c}_{SG}}{dt} = q_p + q_{QL} + q_{GG}, \tag{47}
\]
where
\[
q_p = \frac{1}{a} \frac{3\lambda}{1 - \lambda} \left( -\frac{j_{GL}}{1 + X_p/X_{GL}} \right), \tag{48}
\]
\[
q_{QL} = \frac{D_{SL}c_{SL}}{X_{GL}H_{\lambda}} \left( 1 + \frac{X_{\lambda}}{X_{GL}} \right), \tag{49}
\]
\[
q_{GG} = \frac{1}{a} \frac{3\lambda}{1 - \lambda} \left( -\frac{j_{GL}}{1 + X_p/X_{GL}} \right). \tag{50}
\]
\[ q_{\text{gc}} = -k_{\text{gc}} \tau_{\text{gc}}^2. \]  

(50)

After formal integration,  
\[ \tau_{\text{gc}} = Q_{p} + Q_{\text{gc}} + Q_{Q'}, \]  

(51)

where  
\[ Q_i = \int_0^t q_i(t')dt'. \]  

(52)

4. Numerical solution of generator parameters

Differential equations described in the chapter 3 were solved numerically for general parameters listed in Table 3.

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Label</th>
<th>Magnitude</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diffusion coefficient of chlorine in BHP</td>
<td>D_{CL}</td>
<td>5 \times 10^{-10} \text{ m}^2 \text{ s}^{-1}</td>
</tr>
<tr>
<td>Diffusion coefficient of HO_2^- in BHP</td>
<td>D_{HL}</td>
<td>5 \times 10^{-10} \text{ m}^2 \text{ s}^{-1}</td>
</tr>
<tr>
<td>Diffusion coefficient of oxygen in BHP</td>
<td>D_{\Delta L}</td>
<td>5 \times 10^{-10} \text{ m}^2 \text{ s}^{-1}</td>
</tr>
<tr>
<td>Rate constant of reaction Cl_2 + HO_2^-</td>
<td>k_p</td>
<td>5 \times 10^8 \text{ m}^3 \text{ kmol}^{-1} \text{ s}^{-1}</td>
</tr>
<tr>
<td>Sticking probability of Cl_2 at BHP surface</td>
<td>\gamma_C</td>
<td>0.0075</td>
</tr>
<tr>
<td>Sticking probability of O_2(1\Delta_g) at BHP surface</td>
<td>\gamma_{\Delta}</td>
<td>0.003</td>
</tr>
<tr>
<td>Henry constant of chlorine</td>
<td>H_C</td>
<td>1</td>
</tr>
<tr>
<td>Henry constant of oxygen</td>
<td>H_O</td>
<td>20</td>
</tr>
<tr>
<td>O_2(1\Delta_g) lifetime in BHP</td>
<td>\tau_{\Delta}</td>
<td>2 \times 10^{-6} \text{ s}</td>
</tr>
<tr>
<td>Rate constant of O_2(1\Delta_g) dimolar reactions in (g)</td>
<td>k_{\Delta}</td>
<td>3.7 \times 10^4 \text{ m}^3 \text{ kmol}^{-1} \text{ s}^{-1}</td>
</tr>
</tbody>
</table>

Two examples of calculated results for various input parameters are shown below.

Example 1

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Label</th>
<th>Magnitude</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gas entering the generator</td>
<td>Cl_2 + He</td>
<td>1 : 4</td>
</tr>
<tr>
<td>Total generator pressure</td>
<td>P_{\text{tot}}</td>
<td>40 \text{ kPa (300 torr)}</td>
</tr>
<tr>
<td>Diffusion coefficient of chlorine in gas</td>
<td>D_{CG}</td>
<td>1.04 \times 10^{-4} \text{ m}^2 \text{ s}^{-1}</td>
</tr>
<tr>
<td>Gas temperature</td>
<td>T</td>
<td>260 K</td>
</tr>
<tr>
<td>BHP droplet radius</td>
<td>a</td>
<td>10 \mu m</td>
</tr>
<tr>
<td>Initial HO_2^- concentration in BHP</td>
<td>c_{HL0}</td>
<td>3 \text{ kmol m}^{-3}</td>
</tr>
<tr>
<td>Ratio of BHP/chlorine utilization</td>
<td>U_{H}/U_C</td>
<td>0.5</td>
</tr>
</tbody>
</table>
Results

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Label</th>
<th>Magnitude</th>
</tr>
</thead>
<tbody>
<tr>
<td>Volume fraction of liquid phase</td>
<td>$\lambda$</td>
<td>0.0049</td>
</tr>
<tr>
<td>Radius of gas sphere</td>
<td>$b$</td>
<td>59 $\mu$m</td>
</tr>
<tr>
<td>Initial flux of chlorine per unit droplet surface</td>
<td>$j_{CsI}$</td>
<td>1.7 mol m$^{-2}$ s$^{-1}$</td>
</tr>
<tr>
<td>Initial characteristic depth of O$_2$(1$\Delta_g$) production</td>
<td>$X_P$</td>
<td>0.58 nm</td>
</tr>
<tr>
<td>Characteristic depth of O$_2$(1$\Delta_g$) quenching in liquid</td>
<td>$X_{QL}$</td>
<td>31.6 nm</td>
</tr>
<tr>
<td>Initial flux of O$_2$(1$\Delta_g$)</td>
<td>$j_{\Delta sI}$</td>
<td>1.6 mol m$^{-2}$ s$^{-1}$</td>
</tr>
<tr>
<td>Duration of initial (surface-limited) period</td>
<td>$t_d$</td>
<td>0.24 ms</td>
</tr>
<tr>
<td>Delay of the stage limited by HO$_2^-$ diffusion</td>
<td>$\Delta t$</td>
<td>0.13 ms</td>
</tr>
<tr>
<td>Time of maximum O$_2$(1$\Delta_g$) partial pressure</td>
<td>$t_{\text{max}}$</td>
<td>6.2 ms</td>
</tr>
<tr>
<td>Maximum partial pressure of O$_2$(1$\Delta_g$)</td>
<td>$p_{\text{max}}$</td>
<td>5.5 kPa (41 torr)</td>
</tr>
<tr>
<td>Chlorine utilization</td>
<td>$U_C$</td>
<td>100%</td>
</tr>
<tr>
<td>BHP utilization</td>
<td>$U_H$</td>
<td>50%</td>
</tr>
</tbody>
</table>

**Fig. 4** illustrates a fast HO$_2^-$ depletion from the BHP surface. In this period, the surface process plays a controlling role. Afterwards the overall process is controlled by HO$_2^-$ diffusion. The end of decreasing in HO$_2^-$ mean concentration (at $t \approx 6$ ms) corresponds to exhausting time of chlorine in gas phase. In **Fig. 5**, the first part of the blue curve documents an accumulation of O$_2$(1$\Delta$) during the production period, and the declining part reflects mainly O$_2$(1$\Delta$) quenching in the gas phase. It means that singlet oxygen should exit the generator in a time of the maximum O$_2$(1$\Delta$) concentration, i.e. about 6.2 ms after droplet formation.

![Fig. 4 Time dependence of HO$_2^-$ depletion](image1.png)

![Fig. 5 Time evolution of Cl$_2$ and O$_2$(1$\Delta$) concentration](image2.png)
Example 2

This case was calculated for a higher total pressure (100 kPa (750 torr)), lower BHP concentration, and smaller droplets (d = 2a = 10 µm). The results are in Figs. 6 and 7.

Input data

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Label</th>
<th>Magnitude</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gas entering the generator</td>
<td>Cl₂ + He</td>
<td>1 : 4</td>
</tr>
<tr>
<td>Total generator pressure</td>
<td>Pₜot</td>
<td>100 kPa (750 torr)</td>
</tr>
<tr>
<td>Diffusion coefficient of chlorine in gas</td>
<td>DₜCG</td>
<td>0.33 x 10⁻⁴ m² s⁻¹</td>
</tr>
<tr>
<td>Gas temperature</td>
<td>T</td>
<td>260 K</td>
</tr>
<tr>
<td>BHP droplet radius</td>
<td>a</td>
<td>5 µm</td>
</tr>
<tr>
<td>Initial HO₂⁻ concentration in BHP</td>
<td>cₜ₀HO₂⁻</td>
<td>2 kmol m⁻³</td>
</tr>
<tr>
<td>Ratio of BHP/chlorine utilization</td>
<td>Uₜ/UₜCl</td>
<td>0.5</td>
</tr>
</tbody>
</table>

Results

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Label</th>
<th>Magnitude</th>
</tr>
</thead>
<tbody>
<tr>
<td>Volume fraction of liquid phase</td>
<td>λ</td>
<td>0.018</td>
</tr>
<tr>
<td>Radius of gas sphere</td>
<td>b</td>
<td>19 µm</td>
</tr>
<tr>
<td>Initial flux of chlorine per unit surface of droplets</td>
<td>JₜClI</td>
<td>4 mol m⁻² s⁻¹</td>
</tr>
<tr>
<td>Initial characteristic depth of O₂⁺(Δg) production</td>
<td>XₜP</td>
<td>0.7 nm</td>
</tr>
<tr>
<td>Characteristic depth of O₂⁺(Δg) quenching in liquid</td>
<td>XₜQL</td>
<td>31.6 nm</td>
</tr>
<tr>
<td>Initial flux of O₂⁺(Δg)</td>
<td>Jₜₖₐₜl</td>
<td>3.8 mol m⁻² s⁻¹</td>
</tr>
<tr>
<td>Duration of initial (surface-limited) period</td>
<td>tₜ</td>
<td>0.02 ms</td>
</tr>
<tr>
<td>Delay of the stage limited by HO₂⁻ diffusion</td>
<td>Δt</td>
<td>0.01 ms</td>
</tr>
<tr>
<td>Time of maximum O₂⁺(Δg) partial pressure</td>
<td>tₜₘₐₓ</td>
<td>1.4 ms</td>
</tr>
<tr>
<td>Oxygen pressure at this time</td>
<td>pₜO₂</td>
<td>20 kPa (150 torr)</td>
</tr>
<tr>
<td>Maximum partial pressure of O₂⁺(Δg)</td>
<td>pₜₜₘₐₓ</td>
<td>15 kPa (112 torr)</td>
</tr>
<tr>
<td>Chlorine utilization</td>
<td>UₜCl</td>
<td>97%</td>
</tr>
<tr>
<td>BHP utilization</td>
<td>UₜH</td>
<td>49%</td>
</tr>
</tbody>
</table>
A comparing these results with results of the Example 1 shows that in this case, the initial flux of O_2(^1\Delta_g) and its maximum partial pressure are significantly higher, and all characteristic times are substantially shorter. A maximum partial pressure of O_2(^1\Delta_g) of 15 kPa (112 torr) should be achieved 1.4 ms from the beginning of chlorine contact with BHP droplets. At this time, the calculated O_2(^1\Delta_g) yield is 75%.

5. Two-phase nozzles for spray formation

General considerations

The preliminary calculations showed that high partial pressures of singlet oxygen (≥ 5.5 kPa (40 Torr)) could be achieved with only very small droplets in formed BHP spray (diameter ≤ 20 µm). According to a survey of commercial nozzles available from world producers, small-droplet sprays of fairly viscous liquid (like BHP at low temperatures (10 – 50 mPas)), could be produced by nozzles atomizing liquid by pressurized gas. The air-blast atomizers have the advantage of thorough g/l mixing and production of fine sprays. In a pre-filming type nozzle, liquid is first spread out in a thin continuous sheet and then subjected to atomizing action of high-velocity gas. Two separate gas flows allow the atomizing gas to impact on both sides of the liquid sheet. The nozzle can produce either a hollow cone spray (for a shorter contact time) or a full cone spray.

The crucial factor for choosing a suitable nozzle for the proposed CentSpraySOG is the ratio of gas/liquid flow rates that ensure a time atomization. To obtain the highest Cl_2 utilization, rather low g/l ratio is needed, but to ensure a fine atomization, the gas flow rate must be as high as possible. To exclude a precipitation of the salt from BHP, which could block the nozzle orifices, the nozzles with external atomizing (mixing gas and liquid phases) are preferred. Further, a choice of the nozzle depends also on requirement for the contact time of both phases, which is determined by the
shape of the spray cone. The contact time is shorter in a hollow cone spray than in a full cone spray. The manufacturers give in their prospectus usually only data on the liquid and gas flow rate for different input pressures of liquid and gas but the data on the droplet size are given only exceptionally (and only for water and air at atmospheric pressure and for selected flow rates only).

From the performed survey of available nozzles, we consider following types of the nozzles for our planned application in the investigation of CentSpraySOG:

**BETE Fog Nozzle, Inc.** (50 Greenfield Street, Greenfield, MA 01301)
The SAM-type nozzles with external mixing were designed for spraying of viscous liquid yielding very fine droplets (no data on droplets diameter are given).

**Tab. 4**
Gas (atomizing + fan) flow rate to liquid flow rate ratio for SAM nozzles

<table>
<thead>
<tr>
<th>Type</th>
<th>$v_{G\text{(atom)}} / v_{L}$</th>
<th>$(v_{G\text{(atom)}} + v_{G\text{(fan)}}) / v_{L}$</th>
<th>$(n_{G\text{(atom)}} + n_{G\text{(fan)}}) / v_{L}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>SAM 05-03</td>
<td>0.55</td>
<td>0.82</td>
<td>13.6</td>
</tr>
<tr>
<td>SAM 06-04</td>
<td>0.27</td>
<td>0.42</td>
<td>6.98</td>
</tr>
<tr>
<td>SAM 07-05</td>
<td>0.166</td>
<td>0.25</td>
<td>4.15</td>
</tr>
</tbody>
</table>

**Düsen-Schlick GmbH** (Hutstrasse 4, 96253 Untersiemau/Coburg, Germany) Nozzles of the series 938, 848, 0/28, and 0/48 can atomize liquids to a finer degree than other two-substance nozzles; models 0/48 and 840 are designed for double air atomization.

**Tab. 5**
Atomizing gas flow rate to liquid flow rate for two-substance nozzles providing finest atomization

<table>
<thead>
<tr>
<th>Type</th>
<th>$v_{G}$, Nm$^3$/h</th>
<th>$v_{L}$, l/min</th>
<th>$n_{G} / v_{L}$, mmol (g)/ml (l)</th>
</tr>
</thead>
<tbody>
<tr>
<td>938, 0/28</td>
<td>11.0</td>
<td>0.9</td>
<td>9.1</td>
</tr>
<tr>
<td>938, 0/28</td>
<td>19.0</td>
<td>1.5</td>
<td>9.4</td>
</tr>
<tr>
<td>0/48</td>
<td>28</td>
<td>0.7</td>
<td>30</td>
</tr>
<tr>
<td>0/48</td>
<td>74</td>
<td>2.5</td>
<td>22</td>
</tr>
<tr>
<td>0/48</td>
<td>88</td>
<td>4</td>
<td>16</td>
</tr>
</tbody>
</table>

Water droplet diameter of 3.5 µm to 27 µm is given for nozzle Model 0/48 at water flow rate of 0.4 l/min, and airflow rate of 74 Nm$^3$/h with a spray cone angle of 120°.
Spraying Systems CZ, s.r.o. (Kamenná 31, 639 20 Brno, Czech Rep.)

Tab. 6
Type 1/8J JAU, Internal Mix., no data on droplet size

<table>
<thead>
<tr>
<th>Type</th>
<th>$v_G$, Nm³/h</th>
<th>$v_L$, l/min</th>
<th>$n_G/v_L$, mmol (g)/ml (l)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SUJ 22DF</td>
<td>3.4</td>
<td>0.52</td>
<td>6.6</td>
</tr>
<tr>
<td>SUJ 22DF</td>
<td>6.2</td>
<td>1.33</td>
<td>3.45</td>
</tr>
</tbody>
</table>

For the spray SOG operation using chlorine-helium mixture in the ratio 1:4, and 2 to 8 M HO₂⁻ in BHP, the ratio $n_G/v_L$ of 2.5 to 10 mmol/ml is required for attaining the BHP utilization of 0.5.

6. Model calculations for chosen nozzles

6.1. Results of parametric estimations

The model calculations for the nozzles SAM-05-03 to 07-05 with external g/l mixing and two gas inlets (for atomization gas and gas controlling the spray-cone angle) were performed. The examples below were calculated for gas and liquid flow rates that assure a fine liquid atomization. The pressure was calculated from the molar flow rate of gas and pumping speed corresponding to either the rotary pump capacity (300 m³/h), or this capacity lowered by 2x, 3x, 4x, and 5x, respectively. For these conditions, time dependences of the surface and mean concentration of HO₂⁻ ions in the drop and the mean concentration of chlorine and singlet oxygen in the gas were calculated. They are shown in Figs. 8 – 11. The relative concentration of HO₂⁻ ions in Figs. 8 – 10 is related to the initial HO₂⁻ concentration in the bulk of BHP, the relative Cl₂ and O₂(¹Δ) concentrations in gas are related to the initial Cl₂ concentration in the gas bulk.

Figs. 8 and 9 illustrate time dependences calculated for the nozzle SAM 06-04 and for two different droplet diameters in the spray. It can be seen that HO₂⁻ ions on the drop surface are quickly exhausted (within 1.5 ms). The decrease in the mean HO₂⁻ concentration in liquid finishes after the chlorine exhausting. Neglecting the effect of limited chlorine transport rate in gas causes a sharp change in the course of Cl₂ concentration near zero. A maximum of O₂(¹Δ) concentration was achieved after 24 and 13 ms, respectively. The O₂(¹Δ) yield was still rather high at this time ($Y = 0.64$ or $0.7$, respectively). The Cl₂ exhausting was several times faster in the finer spray (Fig. 9) where the specific surface area of liquid was 4times higher. The Cl₂ and BHP utilizations were also higher for the spray with smaller drops ($U_C = 0.75$ and $U_H = 0.38$).
Fig. 8 Time dependence of relative mean (1) and surface (2) concentrations of HO$_2^-$ ions for 40 µm drop, mean concentration of Cl$_2$ (3) and O$_2$(^1Δ) (4) in gas, and O$_2$(^1Δ) yield (5); $c_{OH_2^-} = 5$ M, $v_L = 31.7$ ml/s, $n_{Cl_2} = 40$ mmol/s Cl$_2$, Cl$_2$:He = 1:4, $Q_{pump} = 0.016$ m$^3$/s, $P_{tot} = 32$ kPa (240 torr).

Fig. 9 Time dependence of relative mean (1) and surface (2) concentrations of HO$_2^-$ ions for 20 µm drops, mean concentration of Cl$_2$ (3) and O$_2$(^1Δ) (4) in gas, and O$_2$(^1Δ) yield (5); other conditions as in Fig. 8.

Results of calculations for a larger nozzle (SAM 07-05) and higher chlorine flow rate (and so a higher pressure), are in Fig. 10.
Fig. 10 Time dependence of relative mean (1) and surface (2) concentrations of HO$_2^-$ ions for 20 µm drops, the relative mean concentration of Cl$_2$ (3) and O$_2$(1∆) (4) in gas, and O$_2$(1∆) yield (5), $n_{Cl_2} = 60$ mmol/s Cl$_2$, Cl$_2$:He = 1:4, $c_{HO2} = 5$ kmol/m$^3$, $v_L = 58.3$ ml/s, $P_{tot} = 48$ kPa (360 torr)

Fig. 11 Time dependence of relative mean (1) and surface (2) concentrations of HO$_2^-$ ions for 20 µm drops, the relative mean concentration of Cl$_2$ (3) and O$_2$(1∆) (4) in gas, and O$_2$(1∆) yield (5), $n_{Cl_2} = 60$ mmol/s Cl$_2$, Cl$_2$:He = 1:4, $c_{HO2} = 3$ kmol/m$^3$, $v_L = 58.3$ ml/s, $P_{tot} = 48$ kPa (360 torr)
The Cl₂ utilization is higher in this case ($U_C = 0.81$) than in Fig. 9, which is caused mainly by a higher surface area of liquid due to the lower g/l ratio for this nozzle (see Table 4). The BHP utilization is slightly lower ($U_H = 0.34$) than in Fig. 9, but it could be higher for less concentrated BHP (3 M HO₂⁻) (Fig. 11). The chlorine utilization is then however lower.

It should be mentioned that the results of calculations above are rather tentative because of the real drop size of the BHP produced by these nozzles is approximate only, and more reliable information about this parameter must be obtained experimentally. Planned measurements on the drop size will be performed first with a model solution of the same viscosity like BHP (NaOH or glycerin), and then with BHP at the temperature used for the COIL operation.

**Pressure dependence of the spray SOG parameters**

**Example 1**

This calculation was performed for different Cl₂ flow rates and one BHP flow rate using the nozzle SAM 06-04. The results correspond to the time of maximum $O_2(1\Delta)$ partial pressure and are shown in Figs. 12 a, b, c.

![Fig. 12 a](image-url)
A decrease in $U_{Cl_2}$ with increasing pressure can be explained by a higher $O_2(1\Sigma)$ loss in gas at higher pressure. At higher pressure, the maximum concentration of $O_2(1\Sigma)$ is achieved at lower degree of $Cl_2$ conversion to $O_2(1\Sigma)$. Chlorine utilization $U_C$ declines markedly with increasing $n_{Cl_2}$ at constant pumping velocity. This decrease is much stronger than it would be correspond to the pressure effect. The BHP utilization ($U_{H}$) weakly increases with increasing $Cl_2$ flow rate. The $O_2(1\Sigma)$ yield decreases with pressure at $p<25$ kPa (~200 Torr). A negligible effect of pressure on the $O_2(1\Sigma)$ yield observed at pressures above 25 kPa (Fig. 12c) is difficult to explain because every point relates to a maximum
$O_2(^1\Delta)$ concentration, not to either constant reaction time or reactor volume. In this example, the highest partial pressure of oxygen was $4 – 5$ kPa (30 – 37 Torr) at medium $O_2(^1\Delta)$ yield of 0.68.

**Example 2**

This calculation was performed for various dilutions of chlorine with helium with constant chlorine flow rate in the nozzle SAM 06-04. At a lower molar fraction of chlorine, the He flow rate was higher resulting in a higher total pressure, while $U_C$ remained constant. This is due to neglecting mass-transfer constrains in the gas phase applied in this model. The results are presented in Figs. 13 **a, b**.

---

**Fig. 13 a, b** Dependence of chlorine utilization, and $O_2(^1\Delta)$ yield on total pressure; nozzle SAM 06-04, $d = 40 \, \mu m$, $v_L = 31.7 \, ml/s$, $n_{Cl_2} = 40 \, mmol/s \, Cl_2$, $x_{Cl_2} = 0.1 – 0.25$, i.e., $n_{He} = 120 – 360 \, mmol/s$, $c_{HO_2} = 5 \, kmol/m^3$
Example 3

In this calculation, the effect of BHP concentration was examined. The results are shown in Figs. 14 a, b.

Fig. 14 a, b. Dependence of chlorine utilization, and \( \text{O}_2(1\Delta) \) yield on BHP concentration at two different pressures; nozzle SAM 06-04, \( d = 40 \mu \text{m}, v_\text{L} = 31.7 \text{ ml/s}, n_{\text{Cl}_2} = 40 \text{ mmol/s Cl}_2 \).

The increasing \( U_C \) with increasing \( \text{HO}_2^- \) concentration is caused by the effect of \( \text{HO}_2^- \) mass transfer rate controlling the rate of \( \text{O}_2(1\Delta) \) formation in the final stage (see the chap. 3). The \( \text{O}_2(1\Delta) \) yield is nearly independent of the \( \text{HO}_2^- \) concentration.
Example 4

This example is calculated for a larger nozzle (SAM 07-05) making possible higher flow rates of BHP at the same gas flow rate, and so a higher ratio of liquid/gas volume. The results are in Figs. 15 a, b.

![Graph 1](attachment:graph1.png)

![Graph 2](attachment:graph2.png)

**Fig. 15 a, b** Dependence of chlorine utilization, and O$_2$(1$\Delta$) yield on total pressure; nozzle SAM 07-05, $d = 40$ $\mu$m, $v_L = 58.3$ ml/s, $n_{Cl_2} = 30 - 60$ mmol/s Cl$_2$, $x_{Cl_2} = 0.2$, $c_{HO_2^{-}} = 5$ kmol/m$^3$

In comparison with Example 1, a higher $U_C$ can be achieved for the same gas flow rate and pressure. This can be explained by a higher number of droplets and their larger specific surface. The O$_2$ partial pressure of 5 – 7.7 kPa (38 – 58 Torr) with O$_2$(1$\Delta$) yield of 0.65 – 0.8 was calculated.
Example 5
This calculation shows the effect of droplet size in the spray on the Cl₂ utilization and O₂(¹Δ) yield (4 times higher specific surface area of liquid vs. Example 1). The results are in Figs. 16a, b, c.

Fig. 16c illustrates the dependence of effective reactor volume on the pressure at Cl₂ molar flow rates that could be sufficient for 1 kW-class COIL. It can be seen that the reactor volume is relatively small (up to 100 cm³) at pressures of 24 – 48 kPa (180 – 360 Torr). The cylindrical reactor of 5 cm in diameter could be then only 5 cm long. The calculated O₂ partial pressure would be 4.8 – 9 kPa (36 – 67 Torr) at the O₂(¹Δ) yield exceeding 0.8.
Fig. 16 a, b, c  Dependence of chlorine utilization, O$_2$(^1$\Delta$) yield, and reactor volume on total pressure; nozzle SAM 07-05, d = 20 µm, $v_L = 58.3$ ml/s, $n_{Cl_2} = 30 - 60$ mmol/s Cl$_2$, $x_{Cl_2} = 0.2$, $c_{HO_2^-} = 5$ kmol/m$^3$

Limitation of the present model

The model used for the above calculations was based on several assumptions, which may result in differences between its prediction and experiment. They are:

1/ Infinite mass transport rate in the gas phase:
This assumption becomes incorrect when the chlorine utilization reaches 100%. At that time, the transport limitation in gas arises and the real utilization would be slightly lower than predicted.

2/ Perfect “co-flow” of gas and liquid phases
In the real situation, the velocity of gas and liquid droplets can be different, especially during the initial stage. Droplets could also coagulate to some extent. Both these effects can result in a surface renewal and so enhancement of the mass transfer towards and backwards the droplet. This effect causes a faster Cl$_2$ utilization and can thus extend a time for singlet oxygen quenching. Quantitative evaluation of the impact of surface renewal on the generator performance can be estimated only experimentally.

3/ The rate of O$_2$(^1$\Delta$) quenching in the liquid phase
The above calculations included the value of O$_2$(^1$\Delta$) lifetime in BHP of $2 \times 10^6$ s given in the literature. Blauer at al. [1] found by comparison between their model and experiments that the liquid phase quenching could be higher by two orders of magnitude than corresponds to the literature value. Based on this estimation, the O$_2$(^1$\Delta$) yields evaluated from our modeling can be treated as the upper limit values.
4/ Uniform droplet size

The real spray will have a certain distribution in the droplet size. The results on Cl₂ utilization and yield are “correct” for the mean droplet radius included in the model calculations, but the droplet size distribution must be taken into account when the model counts also with the enthalpy balance. The present model does not predict yet the heat release and convection, and consequently the amount of water vapor content in the gas. It is obvious that smaller droplets will evaporate faster and so produce more water vapor.

6.2. Conclusions from the modeling

- The calculations performed for chosen nozzles producing the BHP spray with 40-µm or 20-µm droplets show that O₂(^1Δ) can be generated by the CentSpraySOG of a small inner volume 100 cm³, at the oxygen pressure of 4.8–9 kPa (36–67 Torr), and with the O₂(^1Δ) yield exceeding 0.8. The SOG with these parameters would be satisfactory for driving a 1 kW-class COIL. The above calculations were performed for rather concentrated BHP solutions (with 3–7 M HO₂⁻) because the chosen small two-phase nozzles need a relatively high g/l ratio.

- Even higher O₂(^1Δ) partial pressure (20 kPa (150 torr)) could be achieved employing a spray with smaller droplets, but production of such sprays could be difficult due to the high BHP viscosity at low temperature. This problem could be solved by using higher flow rates of more diluted BHP, which would be advantageous also for the lower temperature increase during the reaction with chlorine. A needed lowering the gas/liquid ratio can be obtained easier by using larger nozzles. Temperature of the two-phase mixture could be lowered also by some cooled liquid (e.g., a cooled diluted H₂O₂ or some non-freezing and non-reactive solution), which would be sprayed from another nozzle downstream of the reaction region. After this cooling, the two-phase mixture would enter the centrifugal separator.

7. Experimental set-up for measuring a drop size and drop velocity

7.1. Methods available for these measurements

Methods employed in a drop size measurement may be grouped into three categories: mechanical, electrical and optical. The optical methods are either droplet imaging (photography, holography) or droplet non-imaging (e.g. Phase Doppler Anemometry). Some of them are briefly described below.
**High-speed photography**

This technique is rather simple utilizing a flashlight or laser pulse to create a high intensity and short-time light source. Duration of the flashlight should be of the order of 1 µs, and the laser pulse of the order of nanoseconds. It can be used also for getting the information about drop velocities. This technique is not very convenient because of time-consuming.

**Light-scattering interferometry**

Drop size and velocity measurement is based on the observation of light scattered by drops flying through the crossover region of two intersecting laser beams. The drop velocity is calculated as

\[ U = \frac{\lambda f_D}{2 \sin(\theta/2)} \]

where \( \lambda \) is the wavelength of laser light, \( f_D \) the Doppler frequency, and \( \theta/2 \) laser beam intersection half-angle. A size information is contained in the relative modulation (visibility) of the scattered signal. Based on the scalar diffraction theory, the relationship between drop size and visibility can be expressed as

\[ \text{Visibility} = 2 \frac{J_1(\pi D/\delta)}{\pi D/\delta} \]

where \( D \) is drop diameter, \( J_1 \) the first-order Bessel function, and \( \delta \) the interference fringe spacing.

**Phase Doppler Particle Analysis**

Theoretical description of the dual-beam light scattering showed that the spatial frequency of the scattered interference fringe pattern at off-axis angle is inversely related to a drop diameter. The method uses pairs of detectors at fixed spacing in the image of interference or fringe pattern. The overall measurement size range is from 1 µm to several thousands µm, and can be used also for dense sprays.

**Malvern Particle Sizer**

The device uses a method based on the Freuenhofer diffraction of parallel beam of the monochromatic light by a moving drop. A diffraction pattern is formed, in which part of the light is diffracted depending on the drop size. This commercial technique is very effective, relatively simple and reliable.

The instrumentation of the optical methods is rather expensive and their use needs some experience. It requires calibration with highly mono-dispersed droplets that are either sprayed from a dilute liquid suspension and subsequently dried or present in liquid suspensions.

We plan to employ initially the high-speed photography for exploration of the spray produced by different nozzles. In the next investigation, some non-imagine optical method would be probably necessary.
7.2. Scheme of device for spray parametric study

The device consists of a cooled liquid reservoir connected by a valve with the spray-nozzle. The liquid will be compressed by nitrogen delivered through tube in the upper part. The nozzle is fed by atomizing gas (N₂ or He) producing a spray in the low-pressure chamber evacuated by rotary pump (300 m³/h). Side mirrors of the chamber for the droplet size investigation by the diagnostic beam are purged by means of nitrogen (or He).

Fig. 20. Scheme of device for spray studies
1 - pressurized N₂ (He)
2 – inlet of tested liquid
3 – tank for tested liquid
4 – nozzle
5 – atomizing N₂
6 – diagnostic chamber
7 – diagnostic window
8 – inlet of purging N₂
9 – diagnostic beam

8. Design of gas/liquid blade separator for CentSpraySOG

8.1. Geometry of channels between next blades in separator

The separator is formed by rotor with radial spiral blades. The centrifugal acceleration of the blades is

\[ a = \omega^2 r \] (53)
where \( \omega \) is angular velocity (rad/s) and \( r \) is radius. The angular velocity is related to a number of revolutions per minute (r.p.m.) as

\[
\omega = 2\pi \text{ r.p.m.} / 60 .
\]  

(54)

The centrifugal acceleration can be decomposed into a component tangential to the blade

\[
a_t = a \cos \alpha ,
\]  

(55)

and a component normal to the blade

\[
a_n = a \sin \alpha .
\]  

(56)

Here \( \alpha \) is angle between the tangent of the blade and the radial coordinate.

It is proposed that the blade shape is spiral with polar angle \( \theta \), being a function of the radial coordinate \( r \)

\[
\theta = \theta_{in} - \tan \alpha_{in} - \arctan (1/\tan \alpha_{in}) + (\rho^2 - 1)^{1/2} + \arctan \left( 1/(\rho^2 - 1)^{1/2} \right)
\]  

(57)

Symbols \( \theta_{in}, \alpha_{in}, \) and \( r_{in} \) relate to respective parameters at the inner radius of the blade.

Here \( \rho \) is a dimensionless radial coordinate

\[
\rho = r/q ,
\]  

(58)

where

\[
q = r_{in} / (1 + \tan^2 \alpha_{in})^{1/2}
\]  

(59)

The angle between the blade tangent and radial coordinate is

\[
\alpha = \arctan (\rho^2 - 1)^{1/2} .
\]  

(60)

The length of the blade contour between radii \( r_{in} \) and \( r \) is

\[
L(r) = (r^2 - r_{in}^2) / 2q .
\]  

(61)

This spiral blade has two remarkable properties:

i) The tangential acceleration is constant along the blade surface

\[
a_t = \omega^2 q ,
\]  

(62)

which ensures a driving of the BHP film with constant force.

ii) The height \( h \) of the channel formed by adjacent blades is constant

\[
h = \Delta \theta r \cos \alpha = \Delta \theta r q
\]  

(63)

where \( \Delta \theta \) is a difference of polar angles of adjacent blades of the same radius.

8.2. Hydrodynamics of flowing film along blades

The initial approximation for the flow of liquid along the blade is made for the assumption that most liquid is separated from the g/l mixture at the entrance into the blade space. The relation for the tangential velocity, \( \nu(x) \) was taken from [3] by replacing the gravitational acceleration \( g \) with the centrifugal acceleration \( a \)

\[
\nu(x) = \left[ (\rho \ a \delta^2 \cos \alpha)/2\mu \right] \left[ 1 - (x/\delta)^2 \right] = \left[ (\rho \ \omega^2 \ q \delta^3)/2\mu \right] \left[ 1 - (x/\delta)^2 \right] ,
\]  

(64)
where \( x \) is the coordinate normal to the blade (ranging from \( 0 \) at the liquid/gas interface to \( \delta \) at the liquid/blade interface), \( \mu \) is dynamic viscosity of the liquid.

Maximum velocity at \( x = 0 \) is

\[
\nu_{\text{max}} = \frac{(\rho \omega^2 q \delta)}{2\mu}.
\]

(65)

Average velocity is

\[
\nu_{\text{av}} = \frac{2}{3} \nu_{\text{max}} = \frac{(\rho \omega^2 q \delta)}{3\mu}.
\]

(66)

Volumetric flow rate per one blade of width \( w \) is

\[
V_i = w \delta \nu_{\text{av}} = \frac{(wp \omega^2 q \delta)}{3\mu}.
\]

(67)

Average retention time \( \tau_{\text{av}} \) is

\[
\tau_{\text{av}} = \frac{L(r)}{\nu_{\text{av}}}.
\]

(68)

Depending on the Reynolds number \( \text{Re} = \frac{4 \delta \nu_{\text{av}} \rho}{\mu} \)

(69)

three regimes of the flow can be recognized:

- \( \text{Re} < 20 \): laminar flow with negligible ripping. Here the analysis is fully valid.
- \( 20 < \text{Re} < 1500 \): laminar flow with pronounced rippling. Here the analysis can be good for approximation.
- \( \text{Re} > 1500 \): turbulent flow. The analysis is not valid.

### 8.3. Settling (disengagement) of droplets

Mean displacement due to the Brownian motion is

\[
x_{\text{rms}} = (2 D_p t)^{1/2}
\]

where the diffusion coefficient of the particle (drop) is given as

\[
D_p = \frac{kT C_c}{3\pi \eta d_p}.
\]

(71)

\( C_c \) is the Cunningham correction factor, expressing the velocity slip between the particle and gas. The terminal settling velocity of a particle is

\[
\nu_{TS} = \frac{\rho_p d_p^2 a_n C_c}{18 \eta}.
\]

(72)

Average distance travelled due to a settling is

\[
x_s = \nu_{TS} t = \frac{\rho_p d_p^2 a_n C_c t}{18 \eta}.
\]

(73)

Efficiently removed are particles for which

\[
x_s > x_{\text{rms}},
\]

(74)

having a diameter

\[
d_p > \left( \frac{kT \eta}{(\rho_p a_n C_c t)} \right)^{1/5}
\]

(75)
8.4. Geometry of liquid separator and parameters of liquid film

Dimensions of the rotor of the centrifugal blade separator are given in Table 7.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rotor inner diameter, (d_{in})</td>
<td>50 mm</td>
</tr>
<tr>
<td>Channels number, (n)</td>
<td>6</td>
</tr>
<tr>
<td>Channel width, (w)</td>
<td>50 mm</td>
</tr>
<tr>
<td>Channel length, (l)</td>
<td>100 mm</td>
</tr>
<tr>
<td>Blade revolutions per minute, (n)</td>
<td>1000 – 16000 r.p.m.</td>
</tr>
</tbody>
</table>

Parameters of liquid film

Some examples of the calculated effect of BHP flow rate on the film thickness, the mean velocity of liquid, and the BHP retention time were presented in Figs. 5 – 7 of the interim Report 001. Table 8 gives the film thickness calculated for rotor parameters from the Table 7, the total BHP flow rate of 58.3 ml/s (assumed for the nozzle SAM 07-05 in the Examples 4,5 in chap. 6.1.), and BHP viscosity of 20 mPas.

<table>
<thead>
<tr>
<th>rpm, (\text{min}^{-1})</th>
<th>1000</th>
<th>2000</th>
<th>4000</th>
<th>8000</th>
<th>16000</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\delta), mm</td>
<td>0.4</td>
<td>0.2</td>
<td>0.12</td>
<td>0.09</td>
<td>0.06</td>
</tr>
</tbody>
</table>

The calculated BHP layer in Table 8 is rather thin obviously due to its high tangential acceleration.

Separation of droplets from the spray

Table 9 gives the calculated minimum diameter of droplets that should be separated in the designed separator in dependence on rotation speed. A height of the free space in the channel for the gas flow is 1 mm obtained by subtracting the film thickness from the total channel height. The calculation was performed for the total gas pressure of 50 kPa (375 Torr).

<table>
<thead>
<tr>
<th>rpm, (\text{min}^{-1})</th>
<th>1000</th>
<th>2000</th>
<th>4000</th>
<th>8000</th>
<th>16000</th>
</tr>
</thead>
<tbody>
<tr>
<td>(d_{p,\text{min}}), (\mu\text{m})</td>
<td>2.8</td>
<td>1.8</td>
<td>1.1</td>
<td>0.67</td>
<td>0.33</td>
</tr>
</tbody>
</table>

It is obvious that even small droplets could be separated very efficiently in the proposed centrifugal separator at revolutions above 4000 r.p.m.
8.5. Calculation of gas pressure loss in the channel

- The mean height of the free space in the channel for gas flow, $h_G = 0.8$ mm
- Total volumetric gas flow rate (60 mmol Cl₂ (O₂)/s and 240 mmol He prim/s) at temperature 260 K, and total pressure 50 kPa (375 torr):
  \[ V_{G,\text{tot}} = nRT/P = 0.3 \ R \ 260/50000 = 1.3 \times 10^{-2} \ m^3/h \]
- Total free cross-section for gas flow in 6 channels ($w = 50$ mm, $h_G = 0.8$ mm):
  \[ A_{G,\text{tot}} = 6 \times 0.05 \times 8 \times 10^{-4} \ m^2 = 2.4 \times 10^{-4} \ m^2 \]
- Gas velocity in the channel
  \[ v_{G,1} = \frac{V_{G,\text{tot}}}{A_{G,\text{tot}}} = 54.2 \ m/s \]
- Reynolds number
  \[ \text{Re}_{G,1} = \frac{v \ d \rho_G}{\mu_G} \]
  where gas density \( \rho_G = \frac{P \ M_{av}}{(RT)} = 50000 \times \frac{0.0096}{(8.314 \times 260)} = 0.222 \ kg/m^3 \)
  molecular weight \( M_{av} = \frac{(60 \times 0.032 + 240 \times 4 \times 10^{-3})}{200} = 9.6 \times 10^{-3} \ kg/mol \)
  gas (O₂ + 4He) viscosity \( \mu_G = 1.87 \times 10^{-5} \ Pa \ s \)
  \[ \text{Re}_{G,1} = \frac{v d \rho_G}{\mu_G} = 54.2 \times 0.8 \times 10^{-3} \times 0.222 / 1.87 \times 10^{-5} = 515 \]
  \[ \Rightarrow \text{laminar flow} \]
- Pressure loss in a channel between blades (laminar flow in a rectangular channel):
  \[ \Delta P_{G,1} = 4 \ v \ l \ \mu / (w \ h_G \ k) \]
  \[ k = 0.3 \ h_G/w = 6 \times 10^{-3} \] (tabled value of the coefficient 0.3 is valid for $h_G/w = 0.02$)

The estimated pressure loss $\Delta P_{G,1} = 4 \times 54.2 \times 0.1 \times 1.87 \times 10^{-5} / (0.05 \times 0.8 \times 10^{-3} \times 6 \times 10^{-3})$

\[ = 1688 \ Pa \approx 1.7 \ kPa \]

is acceptable, being about 3% of the total pressure in the generator.

Calculation of O₂(1Δ) loss in the separator

The deactivation of O₂(1Δ) in the gas phase goes through energy pooling

\[ O_2(1\Delta) + O_2(1\Delta) \rightarrow O_2(1\Delta) + O_2(3\Sigma) \quad k_x = 2.7 \times 10^{-17} \ cm^3/s \quad (76) \]

and self quenching

\[ O_2(1\Delta) + O_2(1\Delta) \rightarrow O_2(3\Sigma) + O_2(3\Sigma) \quad k_y = 1.7 \times 10^{-17} \ cm^3/s \quad (77) \]

The effect of O₂(1Δ) quenching can be expressed as

\[ Y^{-1} = (Y_o)^{-1} + k_{tot} \chi \quad (78) \]

where $Y$ is the O₂(1Δ) yield at the beginning and $Y_o$ the yield at the exit,

\[ k_{tot} = k_x = 2 \ k_y \quad (79) \]

and

\[ \chi = (P_{tot})^2 n_{O2} V/(n_{tot} R T)^2 \quad (80) \]
Data for the separator with 6 gas channels: \( P_{\text{tot}} = 50 \text{ kPa}, \; n_{\text{O}_2} = 0.060 \text{ mol/s}, \; n_{\text{tot}} = 0.3 \text{ mol/s} \)

\[ V = 6 \times 0.1 \times 0.05 \times 8 \times 10^{-4} = 2.4 \times 10^{-5} \text{ m}^3 \text{ (volume)} \]

\[ V_{G,\text{tot}} = 1.3 \times 10^{-2} \text{ m}^3/\text{h} \text{ (volumetric flow rate)}, \; \text{and} \; T = 260 \text{ K} \]

Gas retention time in a channel of the length \( l = 0.1 \text{ m} \)

\[ \tau_{G,l} = l/v_{G,l} = 0.1 / 54.2 \approx 1.84 \text{ ms} \]

The factor \( \chi \) calculated from (80) equals to

\[ \chi = 8.56 \times 10^{-3} \text{ mol s/m}^3 = 5.15 \times 10^{15} \text{ molec s/cm}^3 \]

the product \( k_{\text{tot}} \chi \) is then

\[ k_{\text{tot}} \chi = 0.314 \]

and the \( \text{O}_2(1\Delta) \) yield at the separator exit is

\[ Y = 0.64 \]

This yield value is acceptable, although the \( \text{O}_2(1\Delta) \) loss should be still reduced (the product \( P \tau \) is 0.138 torr s). Either smaller height of the gas channel or a shorter channel length could reduce the \( \text{O}_2(1\Delta) \) loss.

9. Design concept of CentSpraySOG device

A cross-section of the rotor of separator was schematically shown in Fig. 3. The perpendicular cross-section of this separator is given on Fig. 21. The rotor consists of two profiled wheels connected (by welding) by six blades. Channels for gas and BHP flow are formed between these blades of the curved shape, see calculations in chap. 8.2.). The reactor inner space is separated from the outer atmosphere by aerodynamic seals formed by a narrow labyrinth gap between rotor and stator. The rotor is connected with the electromotor by a shaft. This motor on one side and a bearing on the other side are protected by purging with nitrogen or helium. The purging gas, which flow rate is much smaller than a gas flow through the nozzle, streams through the labyrinth seal and exits into the reaction space. The shape of trajectory of droplets is affected by the pressure loss in the channels between rotor blades. The higher is this loss, the more uniform is the g/l flow along the blade width.
Fig. 21 Scheme of CentSpraySOG

Dimensions of the CentSpraySOG for about 1 kW COIL are given in the Table 7. The inner diameter and length of the reactor rotor can be significantly increased for attaining higher flow rates of singlet oxygen. The increase in the rotor length is limited by a need of the comparable droplet delay in the inner space for various trajectories.
Fig. 22 shows schematically a possible arrangement of several modules of CentSpraySOG with the outlet into one supersonic laser COIL nozzle.

![Fig. 22 Multi-module CentSpraySOG; 1- spray nozzle, 2- separator body](image)

10. Conclusions of present study on SprayCentSOG

A summary of the results follows the requirements on the advanced SOG characteristics (Objects and goals, p. 4).

1. “*A high pressure operation with ~ 100 torr O₂ partial pressure*”: By our calculations, this parameter could be fulfilled in the proposed CentSpraySOG operating with a droplet spray containing drops less than 20 µm. The calculated chlorine utilization should be above 0.95 and O₂(1∆) yield above 0.75 (see Example 2 in chap. 4, Example 5 in chap. 6.1).

2. “*A single BHP pass burn down*”: The BHP utilization of 0.34 – 0.38 can be attained for the 5 M solution of HO₂⁻ ions, which is one order of magnitude higher utilization than for the jet SOG. Even higher utilization could be attained using more diluted solutions, which needs however to employ nozzles for higher flow rate of liquid (i.e., a lower ratio of gas/liquid flow rate).
3. “A close coupling of the generator to the nozzle plenum to minimize \( \text{O}_2(1^\Delta) \) losses”:
The CentSpraySOG is designed for a direct coupling to the laser nozzle. Our calculations were focused also on estimation of the optimal reactor and separator volume. Fig. 16c illustrates that these requirements are very restrictive. In the reactor volume of \(~100 \text{ cm}^3\), the \( \text{O}_2(1^\Delta) \) yield exceeding 80% could be achieved at chlorine flow rate of 50 – 60 mmol/s and oxygen pressure of 36 – 67 Torr. The calculated overall \( \text{O}_2(1^\Delta) \) yield at the exit from the separator including the reactor (100 cm\(^3\)) and separator (45 cm\(^3\)) was 64%.

4. “An efficient disengagement of gas/liquid mixture”: By our calculations, this process could be efficiently attained by a centrifugal blade separator designed for the CentSpraySOG. At revolutions over 4000 r.p.m., droplets less than \(~1 \mu\text{m}\) should be removed from the gas stream.

5. “A scalability to be suitable for the airborne and mobile application”: One module of the CentSpraySOG can be scaled-up to increase production rate of \( \text{O}_2(1^\Delta) \) by simultaneous increase in the gas and liquid flow rate, the reactor volume and the rate of generation exhausting. A choice of suitable spray nozzles is very broad, and larger two-phase nozzles can even deliver a higher liquid/gas volume ratio. For the need of airborne application, more modules of the CentSpraySOG could be coupled to one laser nozzle (see Fig. 22).

References


Acknowledgement

The investigators are very grateful to the USAF EOARD for the financial support of this work, and thank Dr. Donald Smith, Program Manager, and Mrs. Beth Wann at the EAORD for their assistance and help with the grant arrangements.

We thank Dr. Gordon D. Hager, the grant supervisor, at the US AFRL/DE at the Kirtland AFB, NM, for his encouragement in our work, and beneficial discussions on the grant tasks.