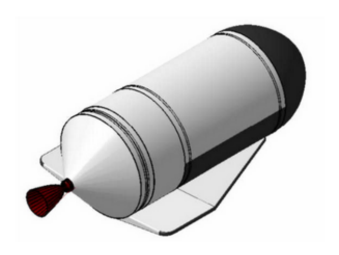


GREDER Green RE-usable DEbris Remover



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Introduction

Ever since the launch of Sputnik 1, the first artificial satellite in October 1957, the number of satellites launched has sharply risen and as of 2020, thousands of satellites are orbiting around the Earth. However, each of every one of those spacecrafts will eventually see their mission being stopped, usually due to a lack of resources from the satellite itself as it reaches the end of its life.

Those who are at the end of their lives will turn into a simple uncontrolled object that keeps orbiting around the Earth and should be avoided as another functioning satellite could take their slot as some of the most important orbits around our planet are starting to get overcrowded and the demand keeps rising.

Moreover, such uncontrolled objects in space can become dangerous as collisions could potentially happen and at such high velocity, those can heavily damage other spacecrafts and create even more debris.

As public awareness grows towards the space debris problem, our mission, Green Debris Remover (GREDER), is looking to contribute to a solution to this problem in a particular orbit, the geo stationary orbit, which is particularly overcrowded due to the many different kinds of satellite operating there.

Schedule

At the start of the project a dedicated group meeting was performed in order to agree on a common project sequence, tasks and challenges as well as work distribution. This group meeting was deemed essential to structure the work packages and to achieve a consolidated baseline for the whole project including time line.

The result is a complex MS project Gantt diagram, which can be found in Annex 1.

1.1 Initial Schedule

The first version of the schedule starts with a project Kick-Off in October which is afterwards followed by a short planning phase. In this planning phase issues as scheduling, work distribution and scope of the project were addressed.

Subsequently, the definition phase started. Within this phase, the vehicle requirements were defined and the mission was planned, calculated and visualized in MATLAB. The outcomes of the definition phase are the boundary requirements which are set to provide a frame for both: the vehicle itself as well as the propulsion system. The requirements were defined at the beginning of the project and were verified after project completion.

Upon definition of the boundary requirements, the specification phase started. Within this phase, different propellant combinations were identified, discussed and compared. Additionally a first mass budget was calculated. The result of this phase is the system specification.

The sequence of the project includes several presentations. The first one was performed in October for a quick overview on the project planning. The second one after the boundary requirements and system specification was set. After this presentation, the vehicle and

sub system design phase started. This phase included major parts of the work packages including propulsion system design with all sub-assemblies as RACS/ACS, propellant tanks, feeding and pressurization system, turbo pumps, catalyzer, engine, injector and nozzle. The outcome of this phase is a preliminary vehicle design and sub system design which was presented in the mid-term presentation.

As a last major work package the simulation phase started. The whole system was simulated including all sub systems and additionally the complex H2O2 decomposition regulation. The final presentation was performed after all tasks were completed and the simulation was finalized.

An overview the compressed initial schedule is shown in Figure 1.1. The detailed Gantt schedule can be found in Annex 1.

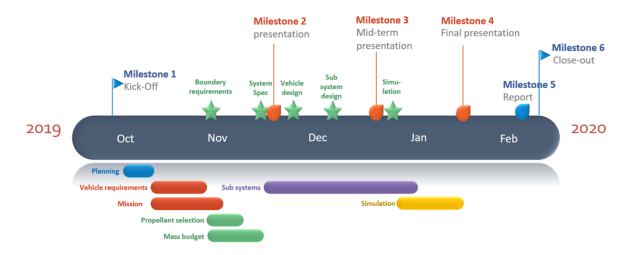


Figure 1.1: Initial schedule

1.2 Final schedule – comparison as planned and as achieved

As usual in project management and project work, not all milestones were achieved in time. As it is shown in the compressed final schedule in Figure 1.2, the finalized vehicle design, the finalized sub-system design and the corresponding simulation shifted within the project schedule (Figure 1.2, shown in red). Nevertheless all work packages have been successfully completed until Milestone 4, the final presentation.

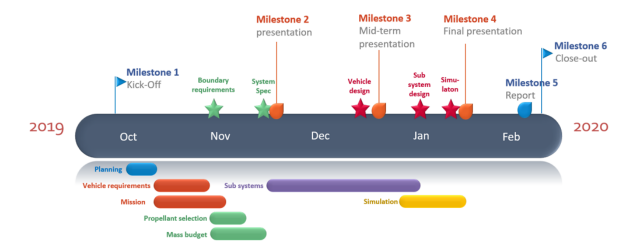


Figure 1.2: Final schedule - comparison as planned and as achieved

Requirements

The top level requirements for the vehicle and propulsion system are divided in three categories: operation, environment and vehicle. The operation requirements are related to the propulsion system and its applicability for the planned missions. The environmental requirements are defined to ensure that both the vehicle and propulsion system is capable of operating and sustaining during launch, mission and in space environment. The scope of the vehicle requirements is to cover the major parts of the planned mission as refuel-ability, aero braking or accuracy.

Operation

TL-1 Provide sufficient thrust for completion of the mission profile including a safety margin
TL-2 Re-ignitable at least 1000 times

TL-3 Service life time of at least 100 missions or 25 years in orbit

TL-4 Ignition and functional reliability shall be higher than 99,5%

Environment

TL-5 Withstand the launch phase

TL-6 Operate in vacuum

TL-7 Withstand the temperature gradients resulting from areas turned towards or away
TL-8 Sustain space-related radiation throughout it's complete life time

TL-9 Withstand debris impact of under 1cm diameter with a max relative speed of 15 km/s

Vehicle

TL-10 The engine shall be the main propulsion system of a GEO satellite recovery vehicle

TL-11 Refuelable between missions

TL-12 Perform aerobrake maneuvers in Earth's atmosphere.

TL-13 Control flight path in Earth's atmosphere using non-propulsive flight control systems TL-14 Remain within the ARIANE 6/Falcon 9 payload launch capabilities to LEO. TL-15 Remain on it's guided trajectory with less than 0.1% deviation.

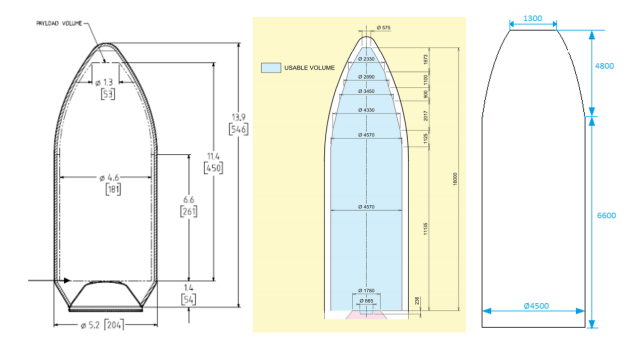
Table 2.1: Requirements for the vehicle

Launcher requirements and launch envelope

From our initial requirements we set to stay within certain margins of size and mass. These two parameters create constraint for the launch. Indeed, we have to analyze the different launcher on the market and their capabilities. Thus, we chose to stay within the capabilities of two of the most used actual launchers: Ariane 6.4 and Falcon 9.

These launchers have almost the same launch capability; the Falcon 9 can send 22.8 t in LEO and Ariane 6.4 can send 21.6 t to LEO. The other important characteristic is the dimension of the fairing. Here we have two different launchers with almost identical size; the Falcon 9 can carry a payload up to 11.4 m long and 4.6 m wide, whereas Ariane 6.4 can carry up to 18 m long and 4.57 m wide but as the Ariane's fairing is very elongated the true usable size will be more around 12 m by 4.57 m.

To allow us to correctly design our spacecraft we need to create a theoretical envelope based on these two launchers. For that we set the following dimension: a diameter of 4.5 m and a total length of 11.4 m. These dimensions are lower than the two launchers to unsure to have a safety margin.



To sum up, here is the 3 fairing we spoke about:

Figure 3.1: Falcon 9 - Ariane 6 - combined fairing

In case, during the design process, the spacecraft cannot stay behind the 22t to LEO launch capability of the Falcon 9, we plan to use the bigger Falcon Heavy which can send 60t to LEO. This launcher will be the last choice, we want at most to stay within our pre-requirement.

We first think to use the structure of our spacecraft, which is aerodynamic, as a fairing so we have less constraint on the size but as it has wings and tail it will generate a small lift but sufficient to destabilize the launcher and might cause a crash. So we decided to forget this idea and to stay within a classic fairing.

Through our project we are going to see if our requirements can meet our calculations and we will make a choice of launcher.

Mission analysis

Satellite catching process

5.1 Choice of the process

In order to catch and de-orbit a satellite in GEO, we considered the following usable tools :

- Net
- Harpoon
- Claw
- Magnet

Other solutions such as towing the satellite or simply removing them from GEO were not considered as they either did not fit our program or would create too much strain on our spacecraft.

We then took a closer look at the feasibility of each solution and compared the advantages and disadvantages :

Solution	${f Advantages}$	Drawbacks	Feasibility		
Net	Cheap, simple, low mass	Slow, hard to handle	Yes (JAXA, ESA)		
Harpoon	Fairly cheap	Can create more debris	Yes (ESA)		
Claw	Safer	Mechanical, moving parts	WIP (CleanSpace One)		
Magnet	Adjustable, no moving parts	Higher mass, needs power	Research State		

As the main focus of our mission is reliability and re-usability, we made the choice of using magnets to catch and hold the satellite we would like to de-orbit.

5.2 Magnetic solution

Even though we decided that we would use magnets, we needed to make sure it was feasible and to lower the drawbacks related to this solution as much as possible. The first precision we need to make is that we will be using electromagnets in order to regulate the intensity of the current in the coil of it, thus, modulating the attraction force so the contact between our spacecraft and the satellite will not be made at high velocity, avoiding damages and space debris creation.

Even though it is still at the state of research, we believe that using electromagnets as our catching solution is realistic as both ESA (with ISAE SupAero) and the NASA have been considering and studying this solution since 2017.

However, as a matter of complexity, we will have to make assumptions in order to simplify the problem. The objective in this part is to prove that, with assumptions, this solution can be applied to our mission and to find the required energy to both catch and hold the satellite until its release.

5.2.1 Assumptions

In our calculations, we assumed that:

- 1. We can consider the magnetic circuit between GREDER and the nozzle of the satellite is a closed one (no air gaps)
- 2. About 0.1% of the satellite's mass is magnetically operable
- 3. Mutual attraction is relatively low compared to the magnetic force
- 4. Residuals in the alloy of the magnets' cores are neglectable

5.3 Sketching and Calculations

Using the first assumption, we can use the formula for the magnetic force in a magnetic circuit with no air gap :

$$F = \frac{(\mu NI)^2 A}{2\mu_0 L^2} \tag{5.1}$$

With:

- μ the magnetic permeability of the core of our magnet (determined by the alloy)
- N the number of turns of the coil around the core

- I the current running through the coil
- A the cross section area of the core
- μ_0 the magnetic constant
- L the length of the mean magnetic circuit

In order to have a good balance between thermal properties and magnetic properties we decided to use an alloy made of 90% of iron and 10% of cobalt for our core. We decided to use this alloy as iron has the best magnetic properties (high relative magnetic permeability) and added cobalt as it has a higher Curie Temperature than iron but has a lower relative magnetic permeability.

In terms of magnet design, we chose to use a squared cross section of $5cm \times 5cm$ for the magnet core and with a length of 15cm, made of an iron-cobalt alloy and a copper coil around it. We decided to have 135 turns of the coil around the core with a coil diameter of 1mm in order to not have the wire revolutions stuck to each other. We also want to run a current of 10A through the coil.

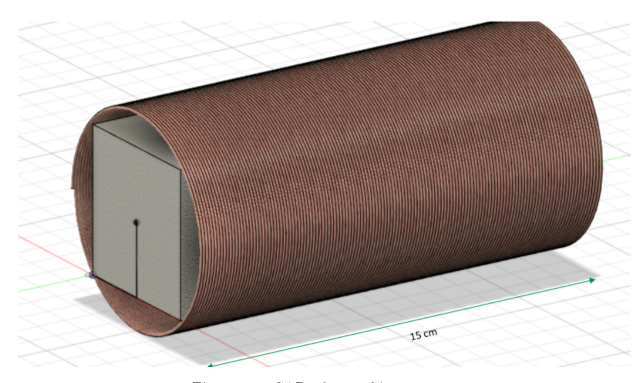


Figure 5.1: CAD of a catching magnet

With those design choices we get the following parameters while taking into consideration that there should not be any kind of residuals in the core alloy:

- $\mu_0 = 4\pi \times 10^{-7} \ H/m$
- $\mu = \left(\frac{\mu_{iron} + \mu_{cobalt}}{2}\right) \times \mu_0 = 5.868 \times 10^{-3} \ H/m$
- N = 135 turns
- I = 10 A
- $A = 25cm^2 = 2.5 \times 10^{-3} m^2$
- $\rho_{core} = \rho_{iron} \times 0.9 + \rho_{cobalt} \times 0.1 = 7.9726 \ g/cm^3$
- $\rho_{coil} = \rho_{copper} = 8.96 \ g/cm^3$

We then need to find the length L of the mean magnetic circuit. In order to do so, we decided to start the catching sequence at 10 m from the satellite and that the target has an exploitable nozzle of 40 cm of diameters which is realistic for spacecrafts in the mass range of 3 500 kg. We could then sketch the catching sequence as so (not to scale):

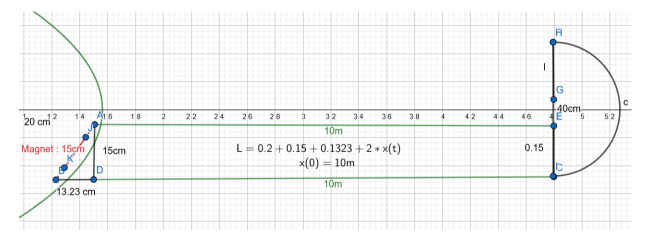


Figure 5.2: Catching sequence (not to scale)

We can then have the length L as a function of the distance between the tip of our spacecraft and the nozzle of our target. As a result we can proceed to find the feasibility of our solution with this magnet design by finding the time it would require at this state to attract the target. However, in this case, the current modulation when the target is close has not been modeled due to its complexity.

Considering that the force will be on one axis only and $m = 0.001 \times m_{target} = 3.5 \ kg$:

$$\vec{F} = m\vec{a} \tag{5.2}$$

$$\frac{(\mu NI)^2 A}{2\mu_0 L(x)^2} = m \times \ddot{x} \tag{5.3}$$

$$\frac{(\mu NI)^2 A}{2\mu_0 [0.2 + 0.15 + 0.1323 + 2x(t)]^2} = m\ddot{x}(t)$$

$$\frac{(\mu NI)^2 A}{2\mu_0 [0.4823 + 2x(t)]^2} = m\ddot{x}(t)$$
(5.4)

$$\frac{(\mu NI)^2 A}{2\mu_0 [0.4823 + 2x(t)]^2} = m\ddot{x}(t)$$
 (5.5)

The catching time can then be found using ode45 on Matlab:

```
clearvars; clc;
    catchtime = 1;
2
   x0 = 10;
3
    while 1
4
        [t,x] = ode45(@f3,[0:1:catchtime], [x0; 0; 0; 0]);
5
            if (x(catchtime ,1) >= 2 * x0)
6
                 break
            else
                 catchtime = catchtime +1;
9
            end
10
   end
11
```

And the function used for the *ode* solver :

```
function [Xdot] = f3(t, X)
mu = 5.686e-3; mu0 = 4 * pi * 10 ^(-7);
N = 135; I = 10; A = 0.05 ^ 2; m = 3500;
x = X(1); y = X(2); vx = X(3); vy = X(4);
Fmag = (mu * N * I) ^2 * A / (2 * mu0 * (2 * norm(X(1:2)) + 0.4823) ^ 2);
Xdot = [vx; vy; 0.001 * Fmag / m; 0];
end
```

We then get a catching time of 812 seconds. As a result, we can determine the energy required to operate the magnets as well as their masses and volumes. We are considering a holding time for approximately half an hour and we also need to verify that the magnets will be able to hold the target while we are de-orbiting.

Heat shield

6.1 Thermal protection system

A complete deorbiting mission from LEO to GEO and returning to LEO has a very high delta-V requirement of roughly $8.5~\rm km/s$. To reduce this delta-V the GREDER spacecraft will use atmospheric drag in the upper atmosphere at an altitude of $70-120~\rm km$ to reduce its relative velocity when returning from GEO to LEO. This kind of maneuver is called an aerobrake and has the potential to save a significant amount of fuel and total spacecraft mass. Aerobrakes are commonly used for reentry vehicles. However, these experience very high thermal loads since the kinetic energy is converted to heat through friction with the atmosphere in a single reentry trajectory. Thus, a high mass for a heat shield is necessary. For a reusable spacecraft ablative heat shields are not useful. Instead, a passively cooled thermal protection system (TPS) was chosen. This allows to radiate the heat away. When the velocity is reduced into small increments the spacecraft can radiate the heat away in the time between the aerobrakes. The advantage is, that fuel is only required for trimming maneuvers to set the targeted perigee radius and not for the decelleration itself. A high number of aerobrakes can be performed with relatively small heat loads if extended mission time is not of a large concern.

Since the GREDER spacecraft is unmanned and does not utilize cryogenic fuels the extended mission time is not as critical. A heat shield needs to be developed for the spacecraft to protect it at the areas with the highest heat loads.

6.2 First estimation

For first rough calculations the total velocity reduced by the aerobrakes was determined as 2.4 km/s. This is the difference in velocity between the perigee velocity of the transfer orbit from GEO to aerobraking altitude and the velocity of the perigee of the transfer orbit

Property	Reinforced Aluminium Carbon-Carbon (2024-T8XX		Steel Type 321	Titanium (6A1-4V)	
Density (kg/m^3)	1578	2803	8027	4437	
Thermal conductivity $(W/m - K)$	3.9*	145.4	14.7	69	
Specific heat $(J/kg - K)$	711.8	816.4	565.2	523.4	
Multipule Use Temperature Limit (K)	1922	425**	872**	662**	
Single Use Temperature Limit (K)	2033	450	922	700	
$T_{cond} \times M_{TempLim}(E+6)$	1.41	0.35	0.49	0.35	

(*thru-the-thickness not isotropic; **values extrapolated by ratio of single/multiple use of CFRC)

Table 6.1: Comparison of materials for heat shield

from the last aerobrake to LEO. The mass of the spacecraft after deorbiting was estimated at 3500 kg. This results in a kinetic energy of:

$$E_{kinetic} = \frac{m_{sc}\Delta v^2}{2} = 10.04 \ GJ \tag{6.1}$$

For a number of 15 aerobrakes this kinetic energy needs to be divided by 15 assuming the temperature at the beginning of each aerobrake will be the same. This kinetic energy was then assumed to be completely converted to thermal energy and distributed evenly over the mass of a heat shield. With the maximum allowable temperature of the chosen material a rough required heat shield mass could be estimated.

6.3Material choice

$$E_{kinetic} = \frac{mv^2}{2}$$

$$m_{shield} = \frac{E_{kinetic}}{c \times dT}$$
(6.2)

$$m_{shield} = \frac{E_{kinetic}}{c \times dT} \tag{6.3}$$

The table above is a comparison of possible TPS materials. The values are taken from the TPSX NASA Material Properties Database. Multiple Use Temperature values were not available for the chosen metals but were extrapolated based on the ratio of the multiple and single use temperature limits of CFRC. The last row shows the product of the values of the lines Specific Heat and Multiple Use Temperature Limit. This relative value allows to quickly compare the materials with euach other regarding the required mass for a heat shield. The unit is J/kg showing that a high value is favorable to store as much energy as possible per mass.

Carbon Fibre Reinforced Carbon quickly proves as the material of choice for a low heat shield mass. Several metals were also taken into account since the manufacturing and integration into the spacecraft struture would be a lot easier. But the low mass for a CFRC TPS in combination with the low thermal conductivity made this the material of choice. CFRC was utilised in the space shuttle heat shield and was found to be the cause of the columbia catastrophe since it is quite brittle and thus sensitive to impact (Columbia Accident Investigation Board Report 1). But since the GREDER spacecraft will be launched inside a rocket launcher fairing no high velocity impacts are to be expected. The capturing of the satellite on the nose cone will be done at very low relative velocities since the force applied by the magnets is controllable with the applied current and can be carefully adjusted (compare chapter 5).

$$m_{shield_{cc}} = \frac{m_{sc} \times (\Delta v_5 \times 1000)^2}{2n_{ab}c_{cc}(T_{max_{cc}} - T_0)}$$
(6.4)

With:

• Δv_5 : Required Δv

• m_{sc} : Spacecraft mass

• n_{ab} : Number of aerobrakes

• T_{max} : Maximum allowable temperature of CFRC

• T_0 : Temperature at beginning of aerobrakes

6.4 Lift and drag

An aerobrake can also be used to change the inclination of the orbital plane. To achieve this the spacecraft needs to be a lifting body. To increase the lift wings can be attached to increase the projeted area of the spacecraft. This area affects both the lift and drag of the spacecraft.

$$F_l = C_l \times \frac{\rho}{2} \times v_1^2 \times A_h \tag{6.5}$$

$$\Rightarrow F_l = 13.9 \ kN \tag{6.6}$$

$$F_d = C_d \times \frac{\rho}{2} \times v_1^2 \times A_h \tag{6.7}$$

$$\Rightarrow F_d = 3.5 \ kN \tag{6.8}$$

¹https://www.nasa.gov/columbia/home/CAIB_Vol1.html

The angle of attack influences the lift and drag coefficients. Since both forces are perpendicular to each other the resulting force and the angle of the direction are calculated by:

$$F_{res} = \sqrt{F_l^2 + F_d^2} (6.9)$$

$$F_{res} = 14.3 \ kN \tag{6.10}$$

$$\beta_{F_{res}} = \arctan\left(\frac{F_l}{F_d}\right) \times \frac{180}{\pi}$$
 (6.11)

$$\beta_{F_{res}} = 76^{\circ} \tag{6.12}$$

It may be noted that unlike an aircraft the lift vector does not point "upwards" but rather parallel to ground and perpendicular to the orbital plane. Thus, the altitude is not affected by the "lift".

The lift and drag coefficient are typically experimental values. For the drag similar shapes can be used as a reference. For the GREDER spacecraft a value of 0.3 is assumed to be reasonable because of its similarity to the bullet shape (compare Figure 6.2 and fig xafter). The delta wing should have a significantly lower drag coefficient and is therefore not taken into account.

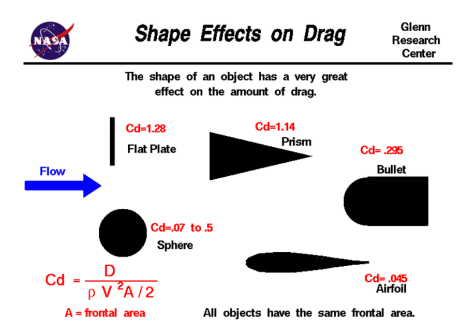


Figure 6.1: Shape effect on drag

Regarding the lift coefficient a strong simplification as made. For a flat plate the lift

coefficient can be approximated with the following formula based on the angle of attack²:

$$C_L = 2\pi \sin(\alpha) \tag{6.13}$$

For an angle of attack of 11° the result for the lift coefficient is 1.2.

To reduce fuel consumption for attitude control a constant angle of attack would be beneficial. To achieve this, the angle between the resulting vector of lift and drag and the velocity vector needs to be the same as the angle between the resulting vector of decelerating and inclination changing vector and the velocity vector. The required delta-Vs for decelerating and inclination change are perpendicular to each other and the angle between them is given by:

$$\beta_{\Delta v_{res}} = \arctan\left(\frac{\Delta v_{inc_{2p}}}{\Delta v_{5}}\right) = 76^{\circ}$$
(6.14)

$$\beta_{F_{res}} = \arctan\left(\frac{C_L}{C_D}\right) \tag{6.15}$$

To compute the angle of attack these formulas need to be equated and the formula for C_L included.

$$\alpha = \arcsin(\frac{\Delta v_{inc2_p}}{\Delta v_5} \times \frac{C_D}{2\pi}) \tag{6.16}$$

For an angle of attack of 11° the angle of the resulting vector of lift and drag is 76°. This results in a balance of both forces during each aerobrake.

6.5 Aerobraking altitude

To estimate the altitude of the aerobrakes, the Matlab function atmoscoesa was used. Here the density of the atmosphere is calculated by altitude between 0 and roughly 84000 m. The values for higher altitudes are extrapolated. A higher altitude results in a longer necessary time in the atmosphere. This has the benefit of a slower heating up of the spacecraft and low values for lift and drag forces. However this could require a higher number of aerobrakes. Weighing these parameters against each other an altitude of 82 km was chosen for the aerobrakes. At this altitude the total aerobraking time is 30.4 minutes. At a number of 20 aerobrakes each would take roughly 91 seconds. During the first aerobrake the distance traveled would be roughly 937 km and during the last roughly 730 km. The forces acted upon the spacecraft are 13.9 kN for lift and 3.5 kN for drag.

²http://brennen.caltech.edu/fluidbook/externalflows/lift/flatplateairfoil.pdf

At a remaining scpacecraft mass of 3000 kg this adds up to an acceleration of 4.8 km/s^2 so roughly 0.5 Gs. These values seemed reasonable since this would keep the mechanical stress relatively low allowing for reusability of the GREDER spacecraft.

6.6 Simplifications

Regarding the thermal loads on the spacecraft the heat shield needs a little more detail. Modeling the heat flux and conductance within the heat shield is simplified due to complexity. Since the even distribution of the heat over the shield is inaccurate several conservative assumptions shall leave a margin for this. CFRC retains its mechanical properties up to 2400° C³. The maximum allowable temperature used for the heat shield modeling is 2000 Kelvin. Also, the assumed temperature at the beginning of the aerorakes is 400 Kelvin. Furthermore It is assumed that the entire kinetic energy is converted to thermal energy and this is completely distributed onto the spacecraft. In reality however not only the spacecraft but also the atmosphere would be heated through friction. This is highly dependent on the geometry of the body. Most reentry vehicles use blunt front faces to "push" a protective shock wave heat shield in front of them. This keeps the highest heat loads away from the surface. This affect can be seen in Figure 6.2 below.

³https://www.makeitfrom.com/material-properties/Carbon-Carbon

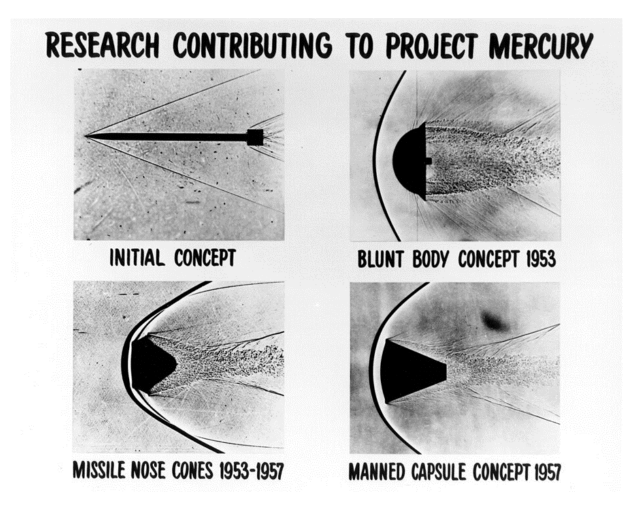


Figure 6.2: Shadowgraph images of Reentry vehicles

As a further simplification, heat transfer is considered solely through convection during the aerobrake maneuver and solely through radiation during the completion of each orbit. The short duration of the aerobrakes is not very significant compared to the duration of a completed orbit (90 s vs. roughly 90 minutes). Furthermore the heat capacity of CFRC increases with increasing temperature. At 750 Kelvin the specific heat capacity is roughly double the value compared to ambient temperature (Specific Heat of Carbon/Carbon Composites ⁴) This leaves an increased margin during the peak heat loads of the atmospheric breaking.

6.7 Heat shield interface

To avoid mechanical failure of the spaceraft's hull beneath the heat shield an insulating blanket will be applied underneath the CFRC heat shield. Herefore, the flight proven

⁴https://apps.dtic.mil/dtic/tr/fulltext/u2/a106709.pdf

AFRSI Blanket was chosen due to its excellent isolating properties and very low density. Here a 10 mm layer of the silica blanket at a density of $\rho_A F = 9.6 \text{ kg/m}^3$ and an additive aereal weight of $m_a dd = 1 \text{ kg/m}^2$ is utilized.⁵

The thermal loads are expected to be highest on the leading edges of the spacecraft's wings and the nose of the spacecraft. In these areas the heat shield shall have the largest thickness. Figure 6.2 shows the areas where the heat shield will be applied. roughly half of the GREDER spacecraft will be covered by the heat shield. This includes the lower faces and leading faces. The upper side and rear of the spacecraft is not expected to endure high thermal loads during the aerobraking maneuvers since the nose will be tilted upwards exposing the lower side.

6.8 Radiative cooling

To verify that the radiative cooling of the CFRC shield is sufficient, an interactive model of the radiated heat was created. To calculate the Power radiated by a gray body in space the following formula was used:

$$P = Aj^* = A\varepsilon\sigma T^4 \tag{6.17}$$

The emissivity of CFRC is $\varepsilon = 0.78$ and σ is the Stefan-Boltzmann constant. ⁶As a starting temperature the maximum allowable temperature of 1982 K was used and for the simulation time the period of the shortest and last orbit with a period of roughly 80 minutes or 4800 seconds. The surface area of the heat shield is half of the total spacecraft surface area.

Calculating the emitted energy could be transferred into a temperature delta for each second of the simulation. after 4800 iterations the temperature is plotted. The last value of 233 K is lower than the assumed 400 K and thus leaves some safety margin.

```
sigma=5.6703E-8 %W/m2K4 stefan boltzmann constant
eps_cc=0.78;
T1=zeros(1,4800); %K
T1(1)=1982; %K
for k=1:1:length(T1)-1
T_rad=sigma*T1(k)^4*eps_cc*A_sc_tot/2/c_cc/m_tps_cc;
T1(k+1)=T1(k)-T_rad;
end
plot(T1);
```

 $^{^5 {}m https://tpsx.arc.nasa.gov/material.html?matid=12}$

 $^{^6}$ https://tpsx.arc.nasa.gov/material.html?matid=24&units=si

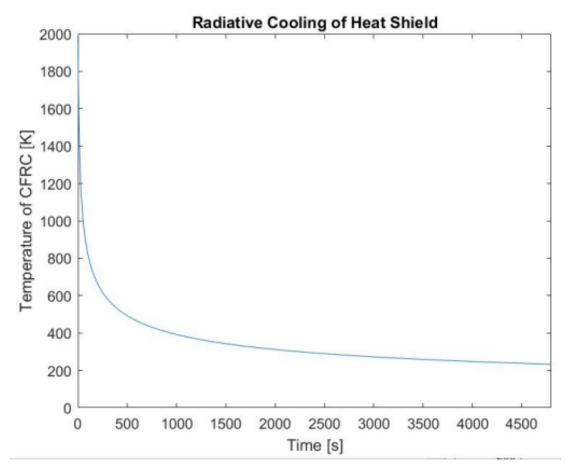


Figure 6.3: Radiative cooling of the heat shield

6.9 Final heat shield mass

For the final mass of the TPS a specific heat capacity for CFRC of $c_{cc}=840~{\rm kJ/kg\text{-}K}$ was assumed. Moreover :

- $\Delta v_5 = 2.396 \text{ km/s}$
- Spacecraft mass $m_{sc} = 3000 \text{ kg}$
- Number of aerobrakes $n_{ab} = 20$
- Starting temperature $T_0 = 400 \text{ K}$

At a surface area of the total spacecraft of $A_{sc_{tot}} = 67.4 \text{ m}^2$ and half of the surface covered with the TPS also the AFRSI blanket mass can be calculated.

$$m_{tps_{cc}} = \frac{(\Delta v_5 \times 1000)^2 m_{sc}}{2n_{ab}c_{cc}(T_{max_{cc}} - T_0)} = 323.9 \ kg$$

$$m_{blanket} = \frac{\rho_{AF}A_{sc_{tot}}}{2 * 0.01} + \frac{A_{sc_{tot}}}{m_{add}} = 37 \ kg$$

$$(6.19)$$

$$m_{blanket} = \frac{\rho_{AF} A_{sctot}}{2 * 0.01} + \frac{A_{sctot}}{m_{odd}} = 37 \ kg \tag{6.19}$$

$$m_{tps} = m_{blanket} + m_{tps_{cc}} = 360.9 \ kg$$
 (6.20)

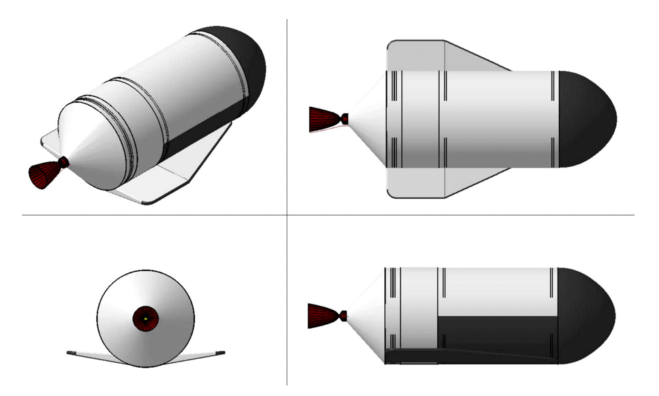


Figure 6.4: GREDER Spacecraft (various views) - Heat shield sown in gray

Propellant selection

For the propellant selection for our vehicle, several important aspects have to be taken into account including:

- Specific impulse in vacuum (Isp) of the propellant
- Storability
- Toxicity including ground handling
- Costs (while the main cost driver is not the propellant itself more its toxicity)
- Reasonable refueling options due to the desired mission profile of the vehicle
- Space flight heritage of the propellant (e.g. flight proven, ground proven or in development)
- Density specific impulse
- Corrosive behavior and compatibility with typical light weight tank materials as titanium or aluminum
- and many more

At first several possible and typical flight proven propellant combinations were analyzed and the most promising combinations were selected and then compared in detail. Afterwards a feasibility study for the chosen propellant combination was deemed necessary in order to check whether the desired propellant combination is able to perform the required mission and de-risk the next development steps.

7.1 Options overview

In literature and historical research several reasonable and flight proven propellant combinations have been found. The propellant combinations are divided in three subcategories: petroleum, cryogenics and hypergolic. These combinations are all bipropellant combinations, the monopropellants have already been excluded at the beginning due considerably low specific impulse in comparison to bipropellants and therefore not suitable for the intended mission profile.

Petroleum fuels are containing a combination of complex hydrocarbons and have been refined from mineral oil. The typical petroleum used as rocket fuel is a highly refined kerosene, called RP-1 (rocket propellant 1). In bipropellant use an oxidizer is necessary and therefore petroleum is often mixed with LOX (liquid oxygen). The specific impulse of a petroleum fuel and cryogenic oxidizer combination is higher than for hypergolic propellant combinations but lower than for fully cryogenic options.

The cryogenics are gaseous bipropellants stored at very low temperatures and usually stand out due to their high specific impulse. The most common cryogenics are liquid hydrogen and liquid oxygen which have to be stored at -253° C for LH2 and -183° C for LOX. Recently, cryogenic combinations with liquid methane (LCH4) as fuel are receiving more attention due to availability methane on mars and therefore it might become attractive for future mars missions.

The last group in this option overview are the hypergolics. The hypergolics are bipropellants that ignite spontaneously when in contact. The main advantage of hypergolics is the storability. Hypergolics are liquid at room temperature and therefore no additional heating or cooling is necessary during the mission. Nevertheless the hypergolics provide less specific impulses than cryogenics and they are mostly highly toxic.

Table 7.1 shows a comparison of different bipropellants divided in the three above mentioned categories. In the initial comparison of this chapters the propellant combinations are compared with regards to their major characteristics: specific impulse, flight evidence and therefore technical risk, storability and their reasonability for our application.

It is clearly visible that most propellant combinations have not been found reasonable for our application due to low temperature storage. Our vehicle is designed to perform several missions with several ignitions permission and refueling at an in orbit refueling station – this mission profile is not conformable with constant low temperature storage.

In contrast to that the group of hypergolics offers great storability and good specific impulse for the mission profile and a storage of these propellants at an in orbit refueling station is feasible. Since the two options MON/MMH and H2O2/RP-1 are promising, further in depth analysis was deemed necessary.

Туре	Combination	Isp [s] vacuum	Flight proven	Storability	Toxicity	Reason- able for our applicati on		
Petroleum	LOX / RP-1	300-353	Used for the lower stages of the Soyuz boosters, and the first stage of the U.S. Saturn V, Atlas, and Falcon 9 boosters. Very similar to Robert Goddard's first rocket.	LOX low temperature storage	non- toxic	NO		
	LOX / LH2	420-453	Used in the stages of the Space Shuttle, Space Launch System, Ariane 5, Delta IV, New Shepard, H-IIB, GSLV and Centaur.	low temperature storage	non- toxic	NO		
Cryogenic	LOX / LCH4 (methane)	355	NO	low temperature storage	non- toxic	NO		
	LF2 (flourine) / LH2	470	NO	low temperature storage	highly toxic	NO		
Hypergolic	MON / MMH	321-336	three first stages Proton booster, Indian Vikas engine for PSLV and GSLV rockets, most Chinese boosters, a number of military, orbital and deep space rockets	storable	highly toxic	YES		
H	H2O2 / RP-1	313	British gamma rocket engines (65kN 2nd stage for example)	storable	non- toxic	YES		

Table 7.1: Propellant combinations overview

7.2 Detailed comparison between MON/MMH and H2O2/RP- 1

The hypergolic propellant combination MON/MMH and H2O2/RP-1 have been found reasonable for our mission and application. Table 7.2 shows a detailed comparison of the propellant combinations. Several characteristics were found to be minor and others were found to be major for our application. The major characteristics are the vacuum specific impulse, the tank volume ration, the combined density, the density specific impulse and the toxicity and storability (highlighted in yellow).

Both propellant combinations have a comparable specific impulse. Therefore no favor

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for either MON/MMH or H2O2/RP-1.

The tank volume ratio is better for MON/MMH because the ratio is nearly one to one, therefore both oxidizer and fuel tank could share the same tank design. This would significantly decrease the development costs since no second tank design and second qualification tank is necessary. Also the manufacturing costs would decreased due to more possibilities of batch production. Especially production costs of forged tank hemispheres are a huge cost driver and these would decrease due to non-recurring costs being distributed on a larger number of hemispheres. Also the costs for jigs and tools would decrease. This favors MON/MMH.

The combined density for $\rm H2O2/RP\text{-}1$ is slightly better than for MON/MMH. Nevertheless the combined density is not a significant value without taking the specific impulse into account. Calculating the density specific impulse, which is basically the specific impulse per mass, $\rm H2O2/RP\text{-}1$ exceeds MON/MMH by 12%. In this category $\rm H2O2/RP\text{-}1$ is the winner because it can provide more specific impulse per mass.

Comparing toxicity of MON/MMH to H2O2/RP-1 it can be stated that MON/MMH is highly toxic and has to be handled with extreme care which increases the ground handling costs of this propellant combination. In contrast to that is the toxicity of H2O2/RP-1. This propellant combination is non-toxic and can be considered as "green" propellant. Nevertheless careful ground-handling is also necessary for this combination because it is prone to reaction with trace elements. H2O2/RP-1 succeeds in this category.

The last category is the storability. Since both propellant combinations are storable in liquid/liquid condition both propellants are suitable for the application.

Taking all criteria and results into account H2O2/RP-1 is the preferred propellant combination for our application. The main reasons are:

- comparable specific impulse to MON/MMH
- better density specific impulse
- non-toxic advantage for maintenance at refuel station
- good on-ground handling
- possibility of R&D funds by ESA for green debris remover

The main reason against MON/MMH is the toxicity.

	MON / MMH	H2O2 / RP-1		
Isp vacuum [s]	321-336	335		
state at 20 °C	liquid / liquid	liquid / liquid		
phase transus solid/gaseous	[-52 °C; 87 °C] / [-11 °C; 21 °C]	[-0,4 °C; 142 °C] / [-60 °C; 300 °C]		
Densities [kg/m³]	1450 / 900	1440 / 900		
Viscosity [mPa-s]	0,86 / 0,48	1,19 / 0,8		
Vapor pressure at 20 °C [bar]	0,05 / 1,1	0,003 / 7,5		
Mixture ratio [Parts of Oxidator]	1,6	7		
Tank volumes per 1000kg propellant [I]	424,4 / 427,4 total 851,6	607,6 / 138,9 total 746,5		
Tank volume ratio	1 to 1,007	1 to 0,23		
Combined density [kg/m³]	1240	1370		
Density specific impulse [Kg*s/m³]	409200	458950		
Chamber temperature [K]	3330	2900		
Toxicity	highly toxic	non -toxic		
Storability	storable	storable		
Corrosion	highly corrosive	slightly corrosive		
Compatible materials	for example Titanium, Aluminum	High purity aluminum (Al95), stainless steel (304, 304L, 316, 316L)		
Isp mono propellant mode [s]	250 / N/A	160 / N/A		
Flight proven	3 first stages Proton booster, Indian Vikas engine for PSLV and GSLV rockets, most Chinese boosters, a number of military, orbital and deep space rockets	British gamma rocket engines (65kN 2nd stage for example)		

Table 7.2: Detailed propellant comparison MON/MMH and H2O2/RP-1

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Propellants	Engine name	Engine type	Engine status	Fvac (kN) Vacuum thrust	OF Ox to fuel ratio	I svac (N- s/kg) Specific vacuum impulse	[ms/mp] Ratio stage empty mass (ms) over total propellant mass (mp)	Calculated Δv [km/s]
H2O2(~0.97)- RP1	RD161P	Upper stage, unknown	Development	24.5 to 14.7	5,9	3128	[]	[]
H2O2(~0.83)- RP1	Gamma-2	Upper stage, Pump fed	Flown	68.2	8,23	2599	0.18	4,46
H2O2()-RP1	BA-44	Upper stage, Pressure fed	Development	196 to 98	7,5	2941	0.08	7,43
H2O2()-RP1	BA-810	Upper stage, Pressure fed	Development	3600 to 1800	7,5	2765	0.08	6,98
								required Δv=6,61

Table 7.3: Historical data of flight H2O2/RP-1 engines

7.3 H2O2/RP-1 feasibility check

Since H2O2/RP-1 is not a common propellant combination and not as frequently used space industry as MON/MHH a further feasibility study has to be performed:

- to de-risk the next development steps
- to check whether the desired propellant combination is able to perform the required mission

In 7.3 historical data of flight H2O2/RP-1 engines is analyzed. One of the engines "Gamma-2" was flown, the others were in development. The thrusts of all engines are comparable or higher than foreseen for our application and two engines also operate in comparable delta-v ranges. This leads to the conclusion that a H2O2/RP-1 engine is generally speaking feasible.

7.4. Summary Page 31/81

7.4 Summary

In summary $\rm H2O2/RP\text{-}1$ is a viable non-toxic hypergolic propellant combination with a good density specific impulse. Additionally the propellant combination is supported by historical flight data. Therefore all further steps will be based on this combination.

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Vehicle design

Chapter 9

Mass model

9.1 Mass Budget - First Iteration

Before actually going into our mass budget, we wanted to get a reference idea for the propellant mass so that we would be sure to be able to achieve our Δv . In order to get this, we decided to find a relation between the usable propellant mass and the mass of the rest as a ratio. This is then fixed and will also allow us to know roughly how much propellant we need depending on the dry mass. Let m_{UP} be the mass of usable propellant. Moreover, we would be aiming for a total initial mass of roughly 20 to 25t on our last iteration. This first iteration was done with another magnet design, presented in November which consisted in two large discs of 600 kg each and have then been abandoned for the second iteration.

9.1.1 Coefficients & Masses after steps

Considering that ISP = 295s and annotating $\frac{m_{UP_i}}{m_{total_i}} = K_i$ with i the burn number :

Step	Required Δv in m/s	K_i	Mass after step
1	2802.4	0.620	$0.38 \ m_{initial}$
2	1342.2	0.371	$0.239 m_{initial}$
3	522.9	0.165	$0.200 m_{initial}$
Satellite caught	NA	NA	$0.2m_{initial}+3500$
4	1487.8	0.402	$0.1196m_{initial}+2093$
Satellite release	NA	NA	$0.1196m_{initial}$ - 1407
5	5.3	0.002	$0.1194m_{initial}$ - 1404.186
6	72.4	0.0247	

9.1.2 Global equation between m_{UP} and $m_{initial}$

Step	$\frac{m_{UP}}{m_{initial}}$	Bias due to debris
1	0.620	0
2	0.141	0
3	0.039	0
4	0.0804	1407
5	0.00024	-2.814
6	0.00295	-36.68
TOTAL	0.88359	+1369.506

We then get our general relation between the usable propellant mass and the initial mass

$$m_{prop} = 0.88359 m_{init} + 1369.506$$

And as $m_{initial} = m_{UP} + m_{rest}$:

$$m_{prop} = \frac{1}{0.11641} \left[0.88359 m_{rest} + 1369.506 \right]$$

 m_{rest} includes the dry mass and the propellant required for the ACS.

9.1.3 First iteration of mass budget

Sub systems

Contributor	Mass in kg
$\overline{ ext{EPS}}$	-
Fuel cells	165.6727
H2 for fuel cell (tank included)	10
Cables	20
GNC	5
Batteries	61.3333
Actuators (for flaps)	10
Servos	1
On board computer	5
$\overline{ m Telecommunications}$	10
Thermal control	10
${ m ACS/RCS}$	-
Reaction wheels	106
ACS (without propellant)	36.16
	440.166

Payload

Contributor	Mass in kg
Magnet	1200

Structure

Contributor	Mass in kg
Hull	509
Wing	54
Engine	60
Engine frame	51
Connectors	25
Tanks	350
Heat shield	472
\underline{Total}	1521

Others

Contributor	Mass in kg
Catalyzer	10
Lines	25
ACS including Propellant	672
Non usable propellant (Residuals, transient, etc.)	200
Helium (including tank)	30
\underline{Total}	937

We then get

$$m_{rest} = m_{Subsystems} + m_{Payload} + m_{Structure} + m_{Others} = 4098.166kg$$

Which, with the previously obtained equation:

$$m_{UP} = 42\ 870.926kg$$

As the mixture ratio is MR = 7.07 and $m_{UP} = m_{UF} + m_{UOP}$

$$m_{UsableFuel} = \frac{m_{UP}}{1+MR} = 5 \ 312kg$$

$$m_{UsableOxidizer} = MR \times m_{UsableFuel} = 37 \ 559kg$$

Results

We can sum this first iteration up with the following table:

Contributor	$\mathbf{Mass}\ (\mathrm{kg})$
Structure	1 521
Magnets	1 200
Sub Systems	440.166
Tank Pressurization	30
Engine	60
Catalyzer	10
Lines	25
Dry mass	3 286.166
Non usable propellant	200
ACS/RCS Propellant	142.12
Usable propellant	42 870.926
Total initial mass	46 969.092

Table 9.7: Initial mass budget

This first initial mass is way over what we are targeting and there are many parameters to be refined during the next iteration.

9.2 Mass Budget - Second iteration

After refining multiple parameters and fixing others to get more accurate values, we went into the second iteration of our mass budget. Having our I_{SP} changed also required another iteration in our calculation formula between the usable propellant mass the the rest of the mass.

9.2.1 Coefficients & Masses after steps

Considering that ISP=315s and annotating $\frac{m_{UP_i}}{m_{total_i}}=K_i$ with i the burn number :

Step	Required Δv in m/s	K_i	Mass after step
1	2802.4	0.596	$0.404 \ m_{initial}$
2	1342.2	0.352	$0.261792 m_{initial}$
3	522.9	0.156	$0.221 m_{initial}$
Satellite caught	NA	NA	$0.221 m_{initial}+3500$
4	1487.8	0.382	$0.137m_{initial}+2163$
Satellite release	NA	NA	$0.137m_{initial}$ -1337
5	5.3	0.0017	$0.1368 m_{initial}$ - 1334.73
6	72.4	0.023	

9.2.2 Global equation between m_{UP} and $m_{initial}$

Step	$\frac{m_{UP}}{m_{initial}}$	Bias due to debris
1	0.596	0
2	0.142	0
3	0.041	0
4	0.084	1337
5	0.0002	-2.273
6	0.0032	-30.699
TOTAL	0.8664	+1304.028

This time our equation between those two masses is given by

$$m_{UsableProp} = \frac{1}{0.1336} \left[0.8664 m_{rest} + 1304.028 \right]$$
 (9.1)

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9.2.3 Second iteration of mass budget

As our way of presenting our first iteration of the mass budget didn't seems clear enough to us, we decided to present it in another, more logical way : **Structure**

Contributor	Mass (kg)
Hull	192
Tanks (including non usable propellant)	700
Wings	136
Lines	60
Connectors	16
$H_2 ank$	12
Structure	1 116

Electrical related contributors

Contributor	Mass (kg)
Batteries	241
Fuel cells	202
On Board Computer	5
Cables	20
H_2 for fuel cells	5
Wing actuators	10
Telecommunications	10
GNC	5
Thermal Control	10
Magnets (Payload)	25.65
Electrical related contributors	533.65

ACS and RCS

Contributor	Mass (kg)
Thrusters	36
H_2O_2	90
Reaction wheels	106
ACS & RCS	232

Propulsion

Contributor	Mass (kg)
Engine	93
Turbopumps	25
Pressurization (He)	1.3
Catalyzer	30
Propulsion	149.3

With those tables, we can deduce m_{rest} :

$$m_{rest} = m_{Structure} + m_{Elec} + m_{ACS\&RCS} + m_{Propulsion}$$
 (9.2)

$$m_{rest} = 2\ 030.95kg \tag{9.3}$$

Thus,

$$m_{UsableProp} = \frac{1}{0.1336} \left[0.8664 m_{rest} + 1304.028 \right]$$
 (9.4)

$$m_{UsableProp} = 22 \ 931kg \tag{9.5}$$

$$m_{Fuel} = \frac{m_{UsableProp}}{MR + 1} \tag{9.6}$$

$$m_{Fuel} = 2841.6kg$$
 (9.7)

$$m_{Ox} = m_{UsableProp} - m_{Fuel} (9.8)$$

$$m_{Ox} = 20\ 089.89kg \tag{9.9}$$

$$m_0 = 24 \ 962.41kg \tag{9.10}$$

In this second iteration with a better I_{sp} and refined values for all of the contributors, we have a large improvement as our initial mass decreased drastically.

Data	Value	Unit
Empty raw mass	2 031	kg
Usable propellant	22 931	kg
Total mass	24 962	kg
Flowrate	10	kg/s
Rocket diameter	2	m
$I_{sp_{vacuum}}$	335	s
Thrust $F = \dot{m}I_{sp}g_0$	32863.5	N
Mixture Ratio	7.07	-
Wall thickness	TBA	m
H_2O_2 internal pressure	1.35	bar

Table 9.10: Frozen information

9.3 Frozen information

[H] After our second iteration of the mass budget, we decided to make a list of the fixed values that we will work around in our further design.

9.3.1 Frozen points

- We will do 20 aerobrakes
- We will have a separate tank design
- H_2O_2 will be pressurized by its decomposition
- The decomposition control will be managed by rotation of the spacecraft
- ACS/RCS Layout similar to the Space Shuttle
- H_2O_2 catalyzers separate
- H_2O_2/O_2 separation via thermodynamic properties
- H_2/O_2 will be used in fuel cells to produce energy

9.4 Mass Budget - Final iteration

As the fixed I_{sp} has been refined as well as other parameters, we went into our final iteration of the mass budget with the same process as the two previous ones.

9.4.1 Coefficients & Masses after steps

Considering that ISP = 335s and annotating $\frac{m_{UP_i}}{m_{total_i}} = K_i$ with i the burn number :

Step	Required Δv in m/s	K_i	Mass after step
1	2802.4	0.574	$0.426 \ m_{initial}$
2	1342.2	0.335	$0.283 m_{initial}$
3	522.9	0.148	$0.241m_{initial}$
Satellite caught	NA	NA	$0.221 m_{initial}+3500$
4	1487.8	0.364	$0.153 m_{initial}+2226$
Satellite release	NA	NA	$0.153m_{initial}$ - 1274
5	5.3	0.0016	$0.1528 m_{initial}$ - 1271.96
6	72.4	0.0218	

9.4.2 Global equation between m_{UP} and $m_{initial}$

5	Step	$\frac{m_{UP}}{m_{initial}}$	Bias due to debris
	1	0.574	0
	2	0.143	0
	3	0.042	0
	4	0.088	1274
	5	0.0002	-2.038
	6	0.0033	-27.73
Τ(OTAL	0.8505	+1244.232

Thus,

$$m_{UsablePropellant} = \frac{1}{0.1495} [0.8505 m_{rest} + 1244.232]$$

9.4.3 Detailed contributors

Structure

Contributor	Mass (kg)
Hull	192
Tanks	700
Wings	136
Lines	70
Connectors	16
Brackets	35
H_2 Tanks	12
Structure	1161

Electrical systems

Contributor	Mass (kg)
Batteries	388.66
Fuel cell	202
OBC	10
Cables	57
H_2 for fuel cells	5
Wing actuators	10
Data transmission	10
GNC	10
Thermal control	20
Magnets	25.65
Electrical systems	738.31

The mass of the batteries is given by $m_{Bat}=\frac{E_{pumps}+E_{magnet}}{ED_{Lithium}}$ with a considered energy density of 100Wh/kg

Attitude control

Contributor	Mass (kg)
Thrusters	36
H_2O_2 for the ACS	90
Reaction wheels	106
Attitude control	232

Propulsion

Contributor	Mass (kg)
Engine	93
Turbopumps + Electric motors	170
He for pressurization	1.3
Catalyzer	40.86
Propulsion	305.16

9.4.4 Final mass budget

Contributor	Mass (kg)
Structure	1161
Electrical systems	738.31
Attitude control	232
Heat shield	360
Propulsion	305.16
m_{rest}	2796.5
m_{UP} with a 5% performance window	25 505
m_{Fuel}	3160.5
m_{Ox}	22 345
Initial wet mass	28 302

Table 9.13: Final mass budget

Chapter 10

Propulsion system

10.1 Engine cycle

The spacecraft uses electrically-driven turbo pumps to feed the oxidizer as well as the fuel to the engine. The propellant and oxidizer are each driven out of their tanks at low pressure, where after a turbo pump in each propellant line strain raises their pressures. As the oxidizer strain has a higher mass flow rate and faces a large pressure drop in the catalyzer, the respective pump is also more powerful as a result. The turbo pumps are driven electrically by electric motors which use large batteries for their power intake. These batteries offer enough charge for one maximum burn time of 900 seconds, after which they are re-powered by a fuel cell which runs on hydrogen and the hydrogen peroxide decomposition product oxygen. This will be explained further in section 10.3. In order to demonstrate how exactly the engine cycle is made up, a flow schematic is shown in the following. Firstly, the pressurization system is shown in Figure 11.1.

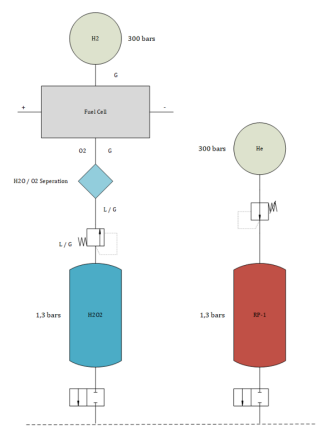


Figure 10.1: Flow Schematic - Pressurization system

As the figure shows, only the RP-1 tank is pressurized by pressurization gas, using a 300 bars helium tank. The hydrogen peroxide has certain decomposition characteristics which enable it to self-pressurize due to the rising pressure upon vaporization. The critical point of hydrogen peroxide is at around 150 degrees Celsius and 1.5 bars, meaning that if the thermal control is sufficiently reliable, a tank pressure of around 1.3 bars can be maintained by self-pressurization. The control system and more details will be explained in section 10.3. The 300 bars H2 Tank that can be seen in the pressurization system flow schematic is therefore not a pressurization tank, but a tank for the sole purpose of running the fuel cell in combination with the oxygen which is separated from water, which is the second decomposition product of hydrogen peroxide. The separation works by simply condensing the water and allowing the gaseous oxygen to pass through a filter. Both the RP-1 tank and the H2O2 tank have a main valve after their outlets, which are visible in Figure 11.1. The remaining feeding system is shown in the second part of the flow schematic, depicted in Figure 11.2.

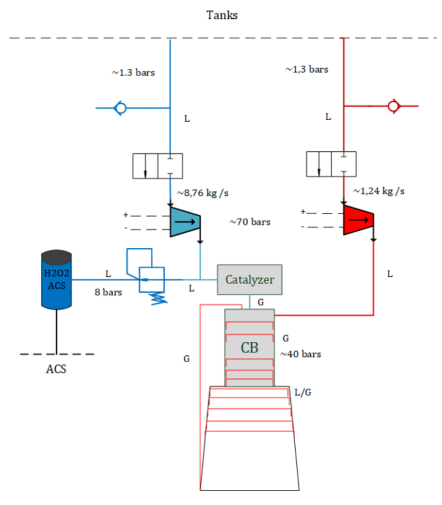


Figure 10.2: Flow Schematic - Engine section

The fueling ports for RP-1 as well as hydrogen peroxide extend to the left- and right-hand-side of the top of the figure. Check valves are situated at these points to only allow propellant flowing in. A second main valve for both propellants is installed just before the turbopumps, which are closed during refueling. While the RP-1 is then funneled through cooling channels in the regenerative cooling system, the oxidizer runs into a catalyzer, where a rapid decomposition reaction splits the hydrogen peroxide into its reactive products for combustion. A second oxidizer strain is guided towards a pressure regulation valve, behind which it continues into a buffer tank of hydrogen peroxide for monopropellant use in the ACS. The catalyzers for ACS thrusters are located in close proximity to their respective combustion chambers. The ACS is not depicted as a flow schematic.

10.2 RCS / ACS

In the previous section we mention that we are going to use hydrogen peroxide as a monopropellant for our Reaction Control System. Indeed, hydrogen peroxide can be used as a quite good RCS propellant.

At the moment, the biggest part of the monopropellant thruster using H_2O_2 are test bench engine. This is because of the difficulty to characterize the engine and its parameters.

Hydrogen peroxide is usable as a monopropellant because of its natural decomposition. As we are going to see on the Catalyzer part further on the report H_2O_2 can be decomposed in H_2 and H_2O . That's this decomposition we are going to use inside our thruster. When hydrogen peroxide goes through a catalyser it decomposes and generates a great amount of heat, up to 1000K. This two factors create steam at a very high temperature and pressure; it's this steam that creates our thrust.

Before designing the engine and its characteristics we need to choose the positioning of the thrusters. In order to do that, we based our design on the American Space Shuttle which use cluster of small thrusters all around the spacecraft to allow a good maneuverability.

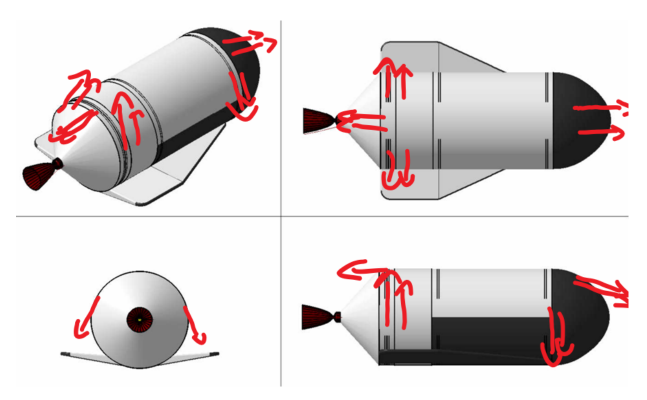


Figure 10.3: RCS thruster repartition

We are going to use 6 different clusters spread over the craft, each cluster is composed of 2 thrusters for a total of 12. This disposition allow to manipulate every axis.

Now we need to compute the characteristic of the thruster, to do so we used RPA (a NASA software to compute the parameters of a thruster based on the propellant and several other characteristics) and some papers of recent studies about hydrogen peroxide thruster.

We assumed a chamber pressure of 10bars and a thrust of 100N, then we use RPA to get the other parameters.

- Combustion temperature: 1223K
- Ejection temperature: 424K
- Ejection pressure : 0.094bars
- $\gamma = 1.335$
- R = 368.6
- ISP = 140s

With these parameters we can then compute every other parameters we want for the thruster and especially the mass flow rate which is necessary to have a correct mass budget.

$$c^* = \sqrt{\frac{RT_c}{\gamma}} \times \frac{\gamma + 1}{2}^{\frac{\gamma + 1}{2(\gamma - 1)}}$$

Throat characteristics:

$$T_t = T_c \left(\frac{2}{\gamma + 1}\right)$$
 $P_t = P_c \left(\frac{2}{\gamma + 1}\right)^{\frac{\gamma}{\gamma - 1}}$ $\rho = \frac{P_t}{RT_t}$ $u_t = \sqrt{\gamma RT_t}$

Exhaust characteristics:

$$M_e = \sqrt{\frac{2}{\gamma - 1} \left[\left(\frac{P_c}{P_e} \right)^{\frac{\gamma - 1}{\gamma}} - 1 \right]} \qquad T_e = \frac{T_c}{1 + \frac{\gamma - 1}{2} M_e^2} \qquad \rho_e = \frac{P_e}{RT_e}$$

Thus we can compute the area ratio and the thrust coefficient:

$$\frac{A_e}{A_t} = \frac{1}{M_e} \left[\frac{2}{\gamma + 1} \left(1 + \frac{\gamma - 1}{2} M_e^2 \right) \right]^{\frac{\gamma + 1}{2(\gamma - 1)}}$$

$$C_F = \gamma \sqrt{\left(\frac{2}{\gamma + 1} \right)^{\frac{\gamma + 1}{\gamma - 1}} \frac{2}{\gamma - 1} \left[1 - \left(\frac{P_e}{P_c} \right)^{\frac{\gamma - 1}{\gamma}} \right]} + \frac{P_e - P_a}{P_c} \frac{A_e}{A_t}$$

Finally we can compute the exhaust velocity, throat area and the mass flow:

$$c = C_F c^*$$
 $A_t = \frac{F}{C_F P_c}$ $\dot{m} = \frac{P_c A_t}{c^*}$

From our assumptions and the results of RPA we finally have:

- Exhaust velocity: c = 617.38m/s
- Mass flow rate: $\dot{m} = 0.105 kg/s$

The most important value for us is the mass flow rate as it allows us to compute the propellant mas needed to perform our maneuvers.

In addition of the thrusters we also need another system which is the Reaction Wheels. This systems which is separated in 3 different wheel, one for each axis, and allow to rotate the entire craft on the 3 axis with the treaction motion applied by the rotation of the wheel.

10.3 Multi-usage of hydrogen peroxide

10.4 Propellant tanks

10.5 Catalyzer

Besides the advantages of H_2O_2 there is one major drawback that we have to take into account. This drawback is that H_2O_2 needs to be decomposed in H_2 and H_2O in order to react with RP-1.

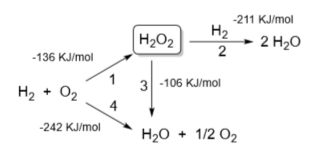


Figure 10.4: H_2O_2 chemical decomposition process

This decomposition is natural but at a low rate whereas we need a very high decomposition rate in order to feed the combustion chamber and sustain a proper flame. This decomposition is an exothermic decomposition. That's why we need a catalyser.

This catalyser needs to be placed between the turbo pump and the injectors. It allows to decompose the H_2O_2 at the last time.

The way a catalyser works is pretty simple; the H_2O_2 goes through a catalyst bed of silver pellets, reacts and generates heat. Why silver? We chose silver because is the mostly used catalyser for H_2O_2 . However, a lot of other different material exists, like Platinum, Manganese or even Gold but these materials are rarely used due to their cost and also the fact that they need to be made in complex alloy in order to optimize the reaction.

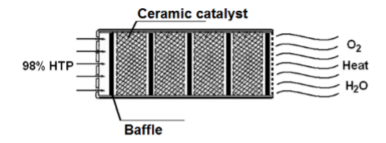


Figure 10.5: Example of catalyst bed

10.5. Catalyzer Page 51/81

The figure above shows basically how a catalyser works. To have an idea of the shape of ours we just have to swap the ceramic catalyst by silver catalyst. So, it will be a steel cylinder filled with small spherical silver pellet separated by some baffles (silver grid mesh).

An important characteristic of the catalyser is the pressure drop it creates. This pressure drop influences the whole feeding system, the turbo pump sizing and even the injector design. that's why we need to characterize the pressure drop created by the catalyser. In order to do so, we are going to use the Ergun equation for packed bed reactor:

$$\frac{\Delta p}{L} = 151.2 \frac{\mu}{d^2} \frac{(1-\epsilon)^2}{\epsilon^2} u + 1.8 \frac{\rho}{d} \frac{1-\epsilon}{\epsilon^3} u^2$$

With μ the dynamic viscosity, ϵ the porosity, d the pellet diameter, ρ the density, L the length of the bed and u the velocity.

In this equation we need some important component such as ϵ . The porosity is complicated to compute and need to be model. According to a recent research we determined a porosity of 0.3802 with a pellet diameter of 5mm.

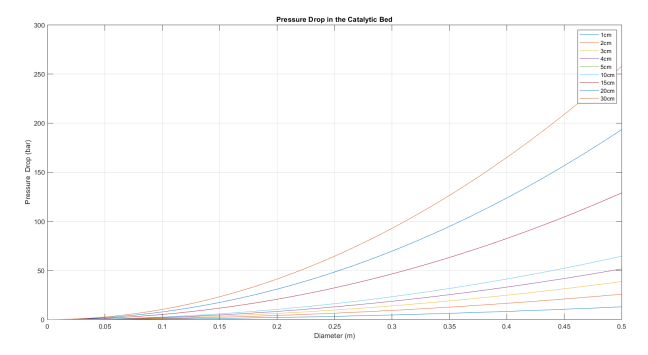


Figure 10.6: Pressure drop depending on the size of the catalyst bed

We see on the graph that the pressure drop rise quickly with the size of the bed. So we need to limit the size of our catalyser in order to not oversize the turbo pump. To do so, we chose to limit our pressure drop to around 30 bars. Finally we obtain a cylinder of 20 cm of diameter and 20 cm of length.

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10.6. Injectors Page 52/81

This geometry allows the catalyser to provide a sufficient decomposition rate, a "contained" pressure drop and size. It also generates a great heat as high as 1000K at the exit of the catalyst bed.

10.6 Injectors

10.7 Feeding system

After having designed most of our propulsion system. We need to carefully link them by designing our feeding system. The biggest challenge is to create a system that will both fit in our spacecraft and deliver the right amount of propellant from the tanks to the engine through our different, required other subsystems.

The general pressure loss in a system is given by:

$$\Delta P = K \frac{\rho}{2} w^2$$

With K depending on the type of change in system geometry as follow:

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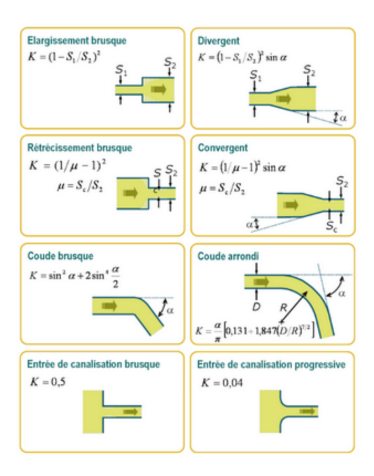


Figure 10.7: K values for geometry changes

In our case, we will use the progressive line entering loss K=0.04 and the bends $K(\alpha)=\sin^2(\alpha)+2\sin^4\left(\frac{\alpha}{2}\right)$. Another K will also be used for the entrance of the injector, which will be specified later on.

For manufacturing costs and simplicity purposes, we choose to only use 45° bends which will result in $K_{bends} = 0.5429$.

With that and the length measurements in mind, we designed the following feeding system layout for which we will then calculate the pressure variations along it:

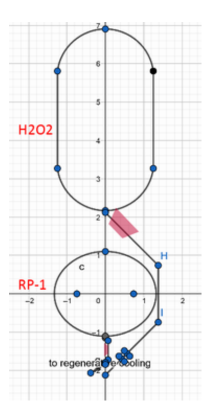


Figure 10.8: Feeding system layout (To scale)

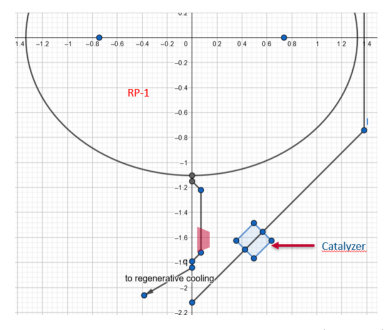


Figure 10.9: Feeding system layout - Zoomed (To scale)

10.7.1 Line diameters

In order to choose our line diameters, we can first use our volume flow and then get the line area from it and then at the end, the line diameter.

Fuel

$$\dot{V}_f = \frac{\dot{m}_f}{\rho_f} = 0.0015298m^3/s \tag{10.1}$$

$$A_{line_f} = \frac{\dot{V}_f}{w_f} = 5.21 \times 10^{-5} mm^2 \tag{10.2}$$

$$d_{line_f} = 2\sqrt{\frac{A_{line_f}}{\pi}} = 8.1447mm \tag{10.3}$$

As we are trying to insure a high injection velocity and to insure a certain margin in pressure and velocity, we will choose a line diameter of 7 mm for the fuel feeding system. Oxidizer

$$\dot{V}_o = \frac{\dot{m}_o}{\rho_o} = 0.006042m^3/s \tag{10.4}$$

$$A_{line_o} = \frac{\dot{V}_o}{w_o} = 0.000275mm^2 \tag{10.5}$$

$$d_{line_o} = 2\sqrt{\frac{A_{line_o}}{\pi}} = 18mm \tag{10.6}$$

In this case, we will choose a line diameter of 15 mm for the oxidizer.

10.7.2 Fuel feeding system

The following items in our fuel feeding system will cause pressure drops:

• Tank exit : K = 0.04

• $4 \times 45^{\circ}$ bends : K = 0.5429 for each

• Straight line losses : $\Delta P = \frac{\rho}{2} w^2 f \frac{L}{D}$

• Friction coefficient : f = 0.02

• Regenerative cooling : $\Delta P = 0.25$ bar

• Fuel injection : $\Delta P = 9.3843$ bars

With our current layout, we have 5 straight lines which will cause pressure losses on the fuel side. Three of them are before the turbopump which is placed at the end of the third straight section, right before the third bend. We consider a velocity of 8 m/s before the turbopumps and of $v_{inj} = 29.363$ m/s after. The line loss for each section is given by :

$$\Delta P = \frac{810}{2} w^2 \times 0.02 \times \frac{L_{section}}{0.007}$$

- 1. First section $(L_{section} = 0.05m)$: $\Delta P = 0.037029$ bar
- 2. Second section $(L_{section} = 0.1m)$: $\Delta P = 0.074057$ bar
- 3. Third section $(L_{section} = 0.5m)$: $\Delta P = 0.37029$ bar
- 4. Fourth section $(L_{section} = 0.1m)$: $\Delta P = 0.997699$ bar
- 5. Fifth section $(L_{section} = 0.05m)$: $\Delta P = 0.49884$ bar

There are also two different values for the bend losses depending on the position of the bend (before or after the turbopump), we have 2 of each:

- $\Delta P_{before} = 0.14072$ bar
- $\Delta P_{after} = 1.8957$ bar

The tank exit loss is:

$$\Delta P_{exit} = K_{exit} \times \frac{\rho_F}{2} \times 8^2 = 0.010368 \text{ bar}$$

10.7.3 Oxidizer feeding system

The following items in our oxidizer feeding system will cause pressure drops:

- Tank exit: K = 0.04
- $4 \times 45^{\circ}$ bends : K = 0.5429 for each
- Straight line losses : $\Delta P = \frac{\rho}{2} w^2 f \frac{L}{D}$
- Friction coefficient : f = 0.02
- Catalyzer : $\Delta P = \text{bars}$
- Oxidizer injection : $\Delta P = 4.9599$ bars

On this part of the feeding system, we also have 5 sections and 4 bends of 45° each. We also consider a velocity of 8 m/s before the turbopump and 21.971 m/s after. However, due to the larger distances (due to our tank layout), the turbopump's position in the feeding system is different and is now positioned after the first bend, right at the beginning of

the second straight line. This results in 3 bends being at high velocity and 1 at relatively slower velocity.

Here, each straight line loss section is given by:

$$\Delta P = \frac{1450}{2} w^2 \times 0.02 \times \frac{L_{section}}{0.015}$$

With:

- 1. First section $(L_{section} = 0.05m)$: $\Delta P = 0.030933$ bar
- 2. Second section $(L_{section} = 1.95m)$: $\Delta P = 9.0992$ bars
- 3. Third section $(L_{section} = 1.4836m)$: $\Delta P = 6.9228$ bars
- 4. Fourth section $(L_{section} = 1.15m)$: $\Delta P = 5.3662$ bars
- 5. Fifth section $(L_{section} = 0.6m)$: $\Delta P = 2.7997$ bars

For the bends, we have:

- $\Delta P_{before} = 0.2519$ bar (1 of them)
- $\Delta P_{after} = 1.0614 \text{ bar (3 of them)}$

The tank exit loss is:

$$\Delta P_{exit} = K_{exit} \times \frac{\rho_o}{2} \times 8^2 = 0.01856 \text{ bar}$$

10.8 Turbo pumps

As most of our subsystems have a defined pressure drop due to their specific design, we have made the choice to use this feeding system design with all losses included to then design our turbopumps to have a pressure rise in accordance with our pressure requirements. We chose to go with electrically driven turbo pumps as we have a good amount of electrical power since we use fuel cells in our spacecraft.

Our respective turbopump required created pressures are:

- Fuel side: $\Delta P_{T_f} = P_{Chamber} + \Delta P_{feeding_f} + \Delta P_{inj_f} + \Delta P_{Regenerative\ cooling} P_{Tank_f}$
- Oxidizer side : $\Delta P_{T_o} = P_{Chamber} + \Delta P_{feeding_o} + \Delta P_{inj_o} + \Delta P_{Catalyzer} P_{Tank_o}$

Thus,

$$\Delta P_{T_f} = 54.395 \text{ bars}$$
 (10.7)

$$\Delta P_{T_0} = 102.28 \text{ bars}$$
 (10.8)

Considering an efficiency of 0.9×0.75 , with 0.9 for the electrical part and 0.75 for the mechanical part, we get the following powers:

$$Power_{fuelpump} = \dot{m}_f \frac{P_{T_f}}{\rho_F \eta} = 13 \ 461 \ W \tag{10.9}$$

$$Power_{Oxpump} = \dot{m_o} \frac{P_{T_o}}{\rho_o \eta} = 96\ 030\ W$$
 (10.10)

Considering a maximum continuous burn time of 900 seconds, we get the following energy with a 40% margin as we are still unsure about the performance of such turbopump:

$$E_{kWh} = \frac{(Power_{fuelpump} + Power_{Oxpump}) \times 900}{3.6 \times 10^6} = 38.322 \ kWh \tag{10.11}$$

We also know the following vapor pressures:

- RP 1: 700 Pa between 20°C and 25°C

With the information we have, we can calculate the pump head rise and the NPSH.

$$H_{p_{Fuel}} = \frac{\Delta p_{p_{Fuel}}}{g_0 \rho_{Fuel}} = 747.474 \ m \tag{10.12}$$

$$H_{p_{Ox}} = \frac{\Delta p_{p_{Ox}}}{g_0 \rho_{Ox}} = 754.192 \ m \tag{10.13}$$

$$NPSH_{Fuel} = \frac{p_{i_{Fuel}} - p_{v_{Fuel}}}{g_0 \rho_{Fuel}} = 6.569 \ m \tag{10.14}$$

$$NPSH_{Ox} = \frac{p_{i_{Ox}} - p_{v_{Ox}}}{g_0 \rho_{Ox}} = 7.0465 \ m \tag{10.15}$$

We can then get the number of stages:

$$n = 1 + floor(\frac{\Delta p_p}{\Delta p_{ps}}) \tag{10.16}$$

Thus, using $\Delta p_{ps} = 47 \times 10^6 \ Pa$,

$$n_{Fuel} = 126\ 373n_{Ox} = 228\ 256 \tag{10.17}$$

We can also get the rotation speeds:

$$N_{Fuel} = 1.636 \ rad/s = 15.623 \ RPM$$
 (10.18)

$$N_{Ox} = 0.532 \ rad/s = 5.079 \ RPM$$
 (10.19)

Then

$$u_{t_{Fuel}} = \sqrt{\frac{gH_{p_{Fuel}}}{n\psi}} = 0.325 \ m/s$$
 (10.20)

$$u_{t_{Ox}} = \sqrt{\frac{gH_{p_{Ox}}}{n\psi}} = 0.243 \ m/s \tag{10.21}$$

$$D_{2t_{Fuel}} = \frac{u_{t_{Fuel}}}{N_{r_{Fuel}}} = 0.199 \ m \tag{10.22}$$

$$D_{2t_{Ox}} = \frac{u_{t_{Ox}}}{N_{r_{Ox}}} 0.457 \ m \tag{10.23}$$

$$D_{1t_{Fuel}} = \sqrt[3]{\frac{\frac{4}{\pi}Q_{Fuel}}{\phi N_{r_{Fuel}}(1 - L^2)}} = 2.197 \ m \tag{10.24}$$

$$D_{1t_{Fuel}} = \sqrt[3]{\frac{\frac{4}{\pi}Q_{Ox}}{\phi N_{r_{Ox}}(1 - L^2)}} = 6.131 \ m \tag{10.25}$$

10.9 Pressure evolution summary

10.9.1 Fuel side

Contributor	Pressure Drop (bars)	Pressure at the end of this part (bars)
Tank	NA	1.3
Tank exit	0.010368	1.29
First section	0.037	1.253
First bend	0.14	1.113
Second section	0.074	1.039
Second bend	0.14	0.899
Third section	0.37	0.529
Turbo pump	54.395 (Rise)	59.924
Valve	5	54.924
Third bend	1.8957	53.0283
Fourth section	0.997	52.0313
Fourth bend	1.8957	50.1356
Fifth section	0.499	49.6366
Cooling	0.25	49.3866
Injection	9.38	40.0066
Combustion chamber	NA	40.0066

Table 10.1: Pressure evolution on fuel side

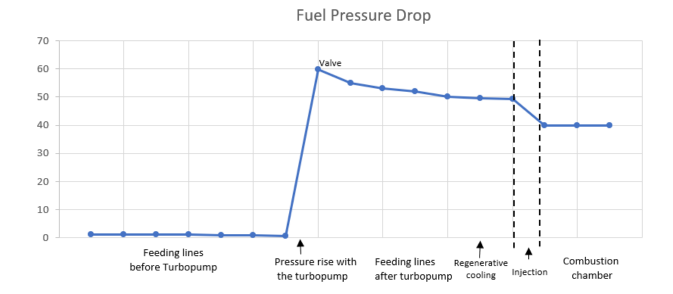


Figure 10.10: Pressure evolution on fuel side (bars)

10.9.2 Oxidizer side

Contributor	Pressure Drop (bars)	Pressure at the end of this part (bars)
Tank	NA	1.3
Tank exit	0.010368	1.29
First section	0.031	1.259
First bend	0.25	1.009
Turbo pump	102.28 (Rise)	108.289
Turbo pump	5	103.289
Second section	9.09	94.199
Second bend	1.06	93.139
Third section	6.9228	86.2162
Third bend	1.06	85.1562
Fourth section	5.366	79.7902
Fourth bend	1.06	78.7302
Fifth section	2.7997	75.9305
Catalyzer	30.95	44.9805
Injection	4.96	40.0205
Combustion chamber	NA	40.0205

Table 10.2: Pressure evolution on fuel side

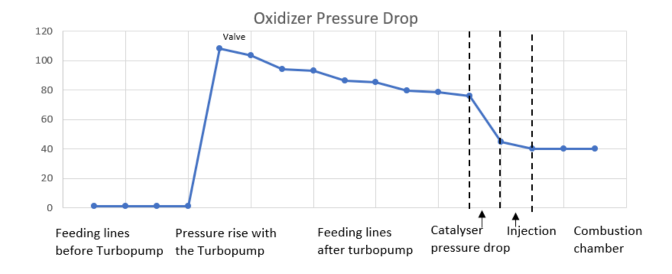


Figure 10.11: Pressure evolution on oxidizer side (bars)

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10.10 Engine

10.11 Nozzle

Chapter 11

Simulations

11.1 Engine performance simulation

In order to simulate the engine performance, the largest single burn was simulated. As the initial burn for entering a Geostationary Transfer Orbit would require more than 900 s of engine burn time due to the large starting mass of the spacecraft, roughly half of this initial burn was simulated. An immediate result of the requirement to limit engine burn time to 900 s is therefore, that the first burn needs to be split into two burns. The engine simulation aims to determine thrust, acceleration, Δv and propellant masses over time. A combined MATLAB® / Simulink® was used, calculating thrust based on following equation:

$$F = \dot{m} \times C_F \times C^* \tag{11.1}$$

with:

$$C^* = \frac{p_c \times A_t}{\dot{m}}$$

and

$$C_F = \sqrt{\frac{2 \times \kappa^2}{\kappa - 1} \times \left(\frac{2}{\kappa + 1}\right) \times \left[1 - \left(\frac{p_e}{p_c}\right)^{\frac{\kappa - 1}{\kappa}}\right]} + \varepsilon \left(\frac{p_e - p_a}{p_c}\right)$$

assuming a constant mass flow rate of $\dot{m}=10~{\rm kg/s}$. Considering a constant chamber pressure after initial start-up of 40 bars and an exit pressure of 3900 Pa, which was determined using RPA, a steady-state thrust of 33.6kN was observed, as can be seen in Figure 11.1. κ was determined using RPA as well and, while varying slightly throughout the engine, assumed as constant to facilitate the simulation.

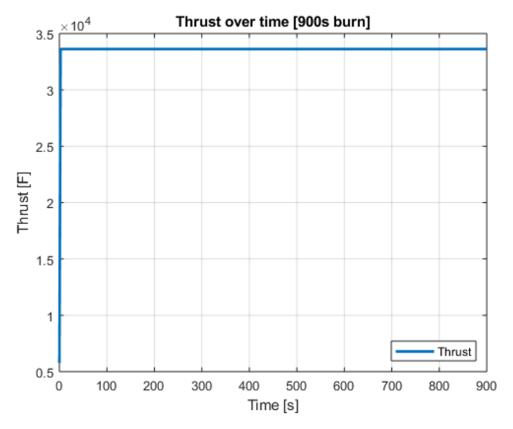


Figure 11.1: Engine simulation - Thrust over time

The resulting acceleration and Δv over time are shown in Figure 11.2.

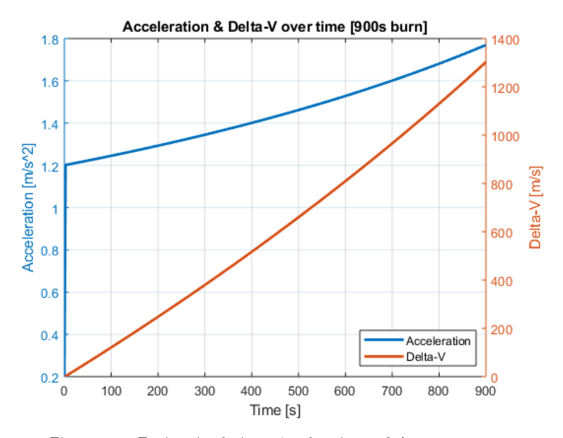


Figure 11.2: Engine simulation - Acceleration and Δv over time

The acceleration, shown in blue, starts off at around 1.2 m/s², which is a value in the range of our expectations. After 900 seconds, the mass loss leads to a final acceleration of slightly below 1.8 m/s², when a Δv of ca. 1 300 m/s is reached. At the point of discovery of the insufficiency of 900 s, the decision to divide the first burn into two was taken. Lastly, the behavior of the propellant masses is portrayed in Figure 11.3.

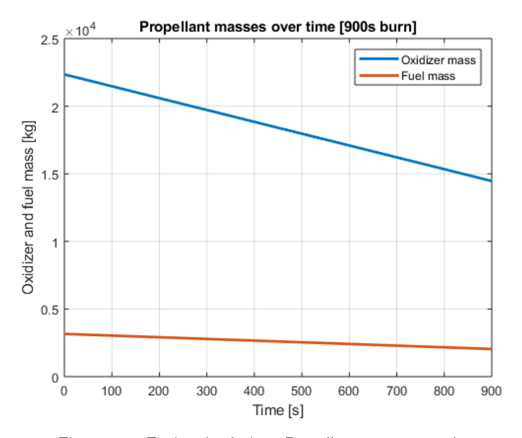


Figure 11.3: Engine simulation - Propellant masses over time

With roughly one third of the total usable propellant having been burnt throughout 900 seconds of burn time, these results are also according to expectations. The simulation of the maximum burn time is representative of the entirety of burns throughout the mission, therefore it validates the viability of the engine design for the fulfillment of mission requirements. The complete simulation script and model can be found in the annex.

11.2 Regenerative cooling simulation

The thermal loads of the engine combustion are managed by regenerative cooling of the engine, with the cooling channels running along axial lines throughout the entire length of the engine. The cooling liquid is the engine fuel (RP-1), being fed into the combustion chamber wall and running along cooling channels with varying cross-section area before exiting at the end of the nozzle. The cooling channel design was determined by simulation of the relevant wall temperatures in steady-state operation, where a temperature at a given location does not change over time, with heat flux in and out being equal. Following assumptions and design choices were made before and during the preliminary calculations:

- The inner engine wall is made of copper throughout the entire engine
- The outer wall is made of steel
- A defined number of symmetrical rectangle-shaped cooling channels run through the engine wall, with varying cross-section to allow for higher and lower thermal flux at different sections
- The cooling channels are symmetrical w.r.t. to the engine propellant flow axis
- Injection into the cooling channels takes place at the combustion chamber end of the engine, in order to create larger heat transfer at more thermally stressed areas due to lower coolant temperature
- The RP-1 is assumed to not change phase during the regenerative cooling (While this decreases simulation accuracy, too little data on high-pressure RP-1 phase change behaviour was available. The temperature does however affect the liquid's heat capacity within the simulation)

In order to simulate the steady-state thermal behaviour of the engine along its length, some simplifications needed to be conducted. All coefficients (e.g. ratio of specific heat capacities) are assumed to be constant throughout each section, with three different values for the combustion chamber, throat area and nozzle respectively. In addition, all time-related values were transformed into distance-based values, so that the simulation time is actually millimetres, instead of seconds. This was achieved by limiting the simulation time to the engine length in millimetres and using switches to change constants based on what section the propellant is in at any given millimetre value, while using the flow velocity as the key parameter to transform into metres. The simulation script, model and complete results can be found in the annex. At this point, only some key conclusions will be presented and discussed.

The central aim of the iterative calculation was to determine a combination of material, wall thickness and cooling channel geometry which allows the inner wall temperature to remain below its maximum operating temperature. All other temperatures, like the coolant

and outer wall temperatures, were of secondary importance. Due to its high thermal conductivity, copper was chosen as the inner wall material.

The cooling channel cross-section was defined to always be equally sized relative to the local engine diameter, with the ratio being a function of the local engine wall cross-section area and a factor, which represents the filling grade. Figure 11.4 shows the cross-section change over the engine length. The rectangular geometry is defined by a ratio of width to height of 5.

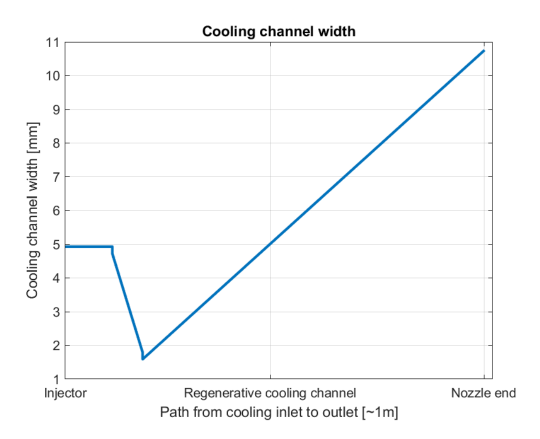


Figure 11.4: Regenerative cooling simulation - Cooling channel width over engine length

The final cooling channel system is a compromise between the copper wall temperature and minimum necessary material for weight savings. Therefore, the copper wall has a thickness of 6 mm in the combustion chamber, 7 mm in the throat area and 5 mm in the nozzle section. The resulting relevant temperatures can be seen in Figure 11.5.

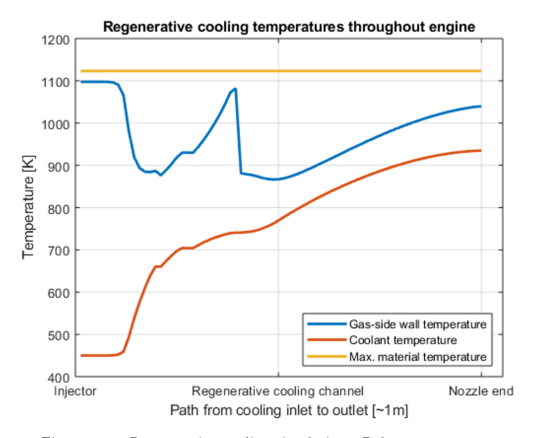


Figure 11.5: Regenerative cooling simulation - Relevant temperatures

The highest temperatures can be observed in the combustion chamber section, as low flow speeds cause less convective heat transfer to take place, reducing the overall heat flux. The gas-side wall temperature then drops in the beginning of the throat area, before rising again towards the throat. After passing the throat, the gas-side wall temperature drops sharply and then rises again towards the nozzle exit.

11.3 Hydrogen peroxide decomposition simulation

As mentioned in section 10.3, the decomposition of hydrogen peroxide is used for self-pressurization of the oxidizer tank as well as power generation in a fuel cell in combination with hydrogen. As hydrogen peroxide is not only decomposing, but the decomposition is also an exothermal process, the thermal control system needs to be well under control in order to avoid critical behavior of the hydrogen peroxide if 150° C are exceeded. Therefore, a detailed simulation has been performed, taking wall thicknesses, materials, radiative and absorbing coefficients as input parameters. The simulation is structured like a control system, calculating the temperature of the hydrogen peroxide inside the tank, the cold and hot wall temperatures. Upon reaching the specified target temperature, the control loop

rotates the spacecraft to a neutral degree for constant hydrogen peroxide temperature. Before discussing the results, an overview of the simplifications taken for facilitation of the simulation follows:

- Inside the tank, only vaporization takes place, no condensation
- The space craft can rotate to any angle instantaneously and there is no modelling of the necessary thruster activation
- The cold and hot walls are always at a homogenous temperature, only depending on heat flux between hydrogen peroxide, the walls, the sun and dark space
- As the structural calculations needed to be complete before the simulation was able to prove the technical feasibility of the decomposition regulation, the hydrogen peroxide tank is assumed to be made of only one material throughout the rest of the documentation, while the simulation allows for two different materials, as different heat flux coefficients are advantageous. Therefore, the results of this simulation are not exactly compatible with the final space craft design and more of a proof of concept, to be reunited with the structural design as a next step.

The simulation showed that, with the simplifications taken, a control system maintaining the hydrogen peroxide pressure at an acceptable level is technically feasible. A control reaction to raising the temperature from 310 K to 350 K has been chosen as the case for technical feasibility demonstration. As explained in section 10.3, the rotational angle of the space craft is the dictating parameter for heat input and output. Figure 11.6 shows the rotational angle, resulting H2O2 pressure, the relevant temperatures and the generated energy considering consumption of all generated oxygen in the fuel cell. The four graphs are outputs of the same simulation, the complete script and model for which can be found in the annex. The initial values for the integration blocks, which output temperatures, are all assumed.

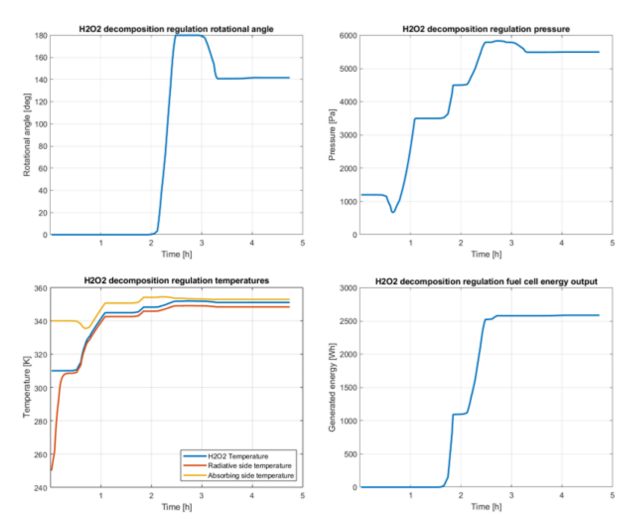


Figure 11.6: Demonstration case for technical feasibility of H_2O_2 decomposition regulation

As the simulation results show, the rotational angle is initially maintained at 0° , meaning full exposure of the more absorbing side towards the sun, with the less radiative side facing dark space. After 2 hours, the space craft turns towards the other side, decreasing the heat flux gradually while approaching the target temperature. In the end, a constant neutral rotational angle of 140° is maintained, with a final control error of around 1-2 K. After this process, 2,5 kWh of energy were generated, while the pressure inside the tank has risen from around 1000 Pa to 5500 Pa. The tank pressure is determined via interpolation of temperature-dependent vaporization pressures to gain a function of pressure with respect to temperature.

Chapter 12

Evaluation

- 12.1 Requirement verification
- 12.2 Lessons learnt

Conclusion

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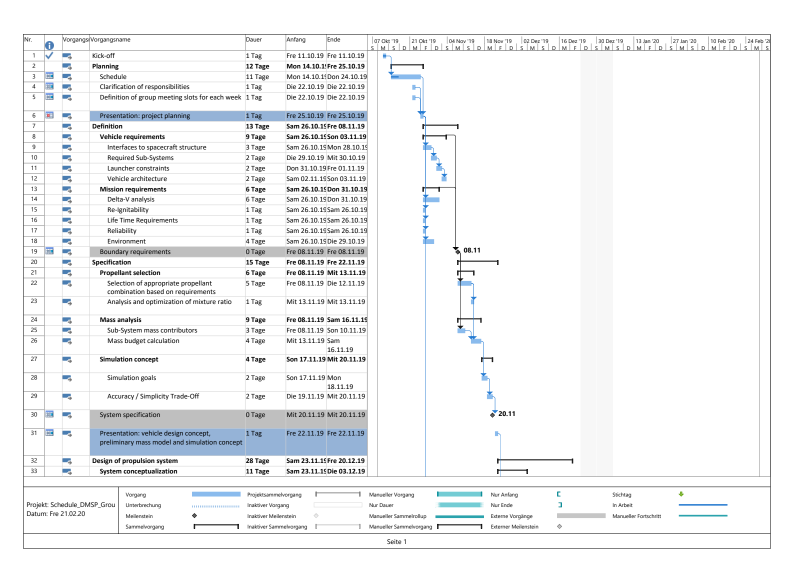
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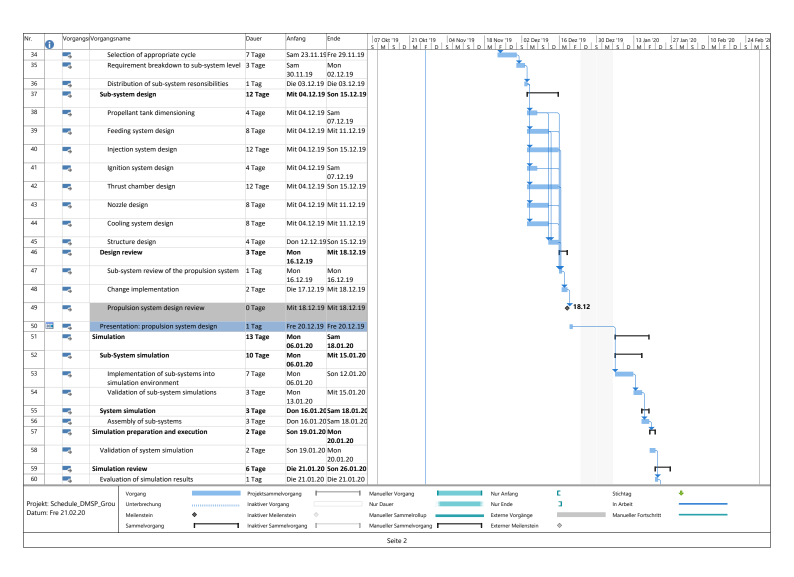
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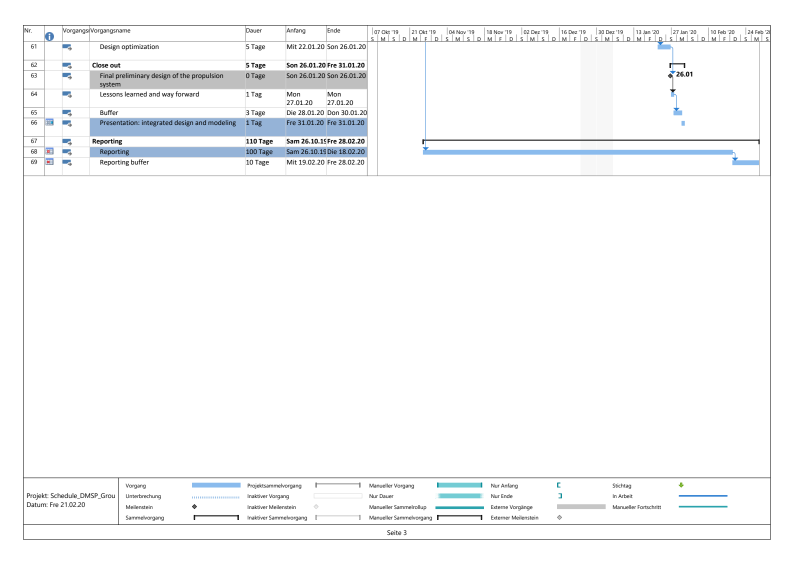
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Annex I - Gantt Diagramm







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