Contamination management pertains to the analysis and optimization of processes with regard to the cleanliness of components, systems and the purity of the fluids used. In today's hydraulic systems, the hydraulics and mobile hydraulics industry, smaller, lighter and more powerful components are currently being used compared to 10 years ago. The use of these components also means that the demand of system cleanliness is now much higher, as has been shown by various studies.

As much as 90% of all hydraulic system outages is due to increased contamination. This failure rate applies to more than classic hydraulics industry. Contamination Management is also a key issue in the automotive industry, in which the use of electrohydraulic systems is on the rise. In this context, hydraulic or fluid power systems are used in a general sense for all industries.

In the past, fluid power systems were equipped with system filtration, which cleaned the system during commissioning and then had the task of maintaining system fluid cleanliness at a constant level, e.g. by using commissioning filters and initial brief maintenance intervals followed by changing over to system filtration. This approach frequently no longer suffices due to the growing demands made of today's hydraulic systems (extended maintenance intervals and mounting cost pressure). Precommissioning flushing is performed in large systems in the hydraulics industry to quickly bring the contamination down to an acceptable level.

However, in small, mass-produced hydraulic systems this is not always possible. That is why contamination management begins with the manufacturing of individual components and extends throughout the entire process up to and including the finished component. Ideally, the design and development departments are integrated in this process so that component design facilitates the washing of components in a cost-efficient manner. Suppliers also have to be involved in contamination management when the manufacturing process involves a large portion of sourced components. By introducing contamination management to minimize particulate concentration in all areas, beginning with manufacturing and extending to the operation of the entire system, system malfunction and failure caused by particulate contamination can be prevented. As a result, costs savings are achieved. Cutting the costs of machining tools, improving the utilization of test stations, and optimizing the use of washing machines can help do this.

This results in the following contamination management tasks: development of systems which are optimized so as to facilitate cleaning, optimizing and monitoring washing and flushing processes, training employees and raising their awareness, detecting and eliminating sources of contamination, drafting analysis instructions, and drafting cleanliness specifications for components and systems. An overall cost assessment is done to gauge the success of contamination management. Consider the following factors: warranty and non-warranty courtesy work, energy costs, reworking costs, operating costs of washing machines and test stations, machining tool costs, and labor costs.

### Definitions

**Contamination Management**
Monitoring/optimization of cleanliness in material flows and system assembly

**Fluid Power System**
Hydraulic systems, including automotive systems containing fluid fillings

**Basic Contamination**
Quantity of contamination present subsequent to assembly

**Ingress Contamination**
Particulate contamination caused by ingestion

**Initial Damage**
Damage to surfaces caused during function testing/commissioning/assembly of systems

**Contamination Monitoring**
Analysis of processes with regard to the ingress of dirt caused by them

**Online Measurement**
Measurement process which the sample to be analyzed is process fed to a measurement device directly from the system

**Offline Measurement**
Measurement process in which the sample is taken from the process system and analyzed elsewhere, e.g. taking an oil sample and sending it in to a laboratory
Various types of contamination occur in fluid power systems: gaseous (e.g. air), liquid (e.g. water) and solid contaminants. An overview of the various contamination types is shown in the following diagram.

As you can tell from examining Figure 1, solid contamination is subdivided into three groups: extremely hard, hard and soft particles. Extremely hard particles can cause substantial damage in fluid power systems if they are not removed as quickly as possible. Preventive measures can reduce the ingress of contaminants in systems.

Figure 1. Types of Contamination

Hard particles are frequently listed separately in specifications. Maximum values are specified for the longest dimension these hard particles may have, e.g. largest abrasive particle: max. 200 µm or 200 x 90 µm or no particles > 200 µm.

Not only the hardness of contamination particles play a role but also their number and size distribution as well.

The particle size distribution in new systems is different from that of systems that have been in operation for a number of hours. In new systems, there is an accumulation of coarse contaminants up to several millimeters long, which are then increasingly reduced in size in the course of operation or eliminated by filtration. After several hours of operation most particles are so small that they are no longer visible to the naked eye.

When commissioning fluid power systems there is additional particulate contamination by virtue of abrasive wear in which rough edges are worn away through running-in. Contamination management can’t prevent this ingress of contaminants; however, if basic contamination is lower, there is less abrasion during system startup.
As Figure 2 shows, the level of contamination without contamination management is higher throughout system operation as compared to a system in which contamination management is employed, the result being that more initial damage may be caused to surfaces.

**Figure 2. Cleaning of a Fluid Power System With and Without Contamination Management**

Microscope images show typical particle samples, containing fine particles, as occur in fluid power systems. (Figure 3)

**Figure 3. Typical Particle Samples**

An average healthy human eye can see items down to approximately 40 µm in size. Particle analyses are conducted using a microscope or in fluid power systems using particle counters employing the light extinction principle. (Figure 4)

**Figure 4. Sizes of Known Particles in Inches and Microns**
Particulate contaminants circulating in fluid power systems cause surface degradation through general mechanical wear (abrasion, erosion, and surface fatigue).

This wear causes increasing numbers of particles to be formed, the result being that wear increases if the "chain reaction of wear" is not properly contained (by reducing contamination).

Gaps grow larger, leakage oil flows increase in size, and operating efficiency (e.g. of pumps) decreases. Metering edges are worn away, thus resulting in control inaccuracies. In some cases, blockage of control ducts or nozzle bores occurs.

The chain reaction of wear during the everyday operation of hydraulic systems has to be interrupted by properly designed and dimensioned filter systems. However, the measure of security afforded the user is deceptive as highly damaging contaminants seep in during component and system assembly and system installation. This ingress of contaminants not only can cause preliminary damage to system components but also premature failure as well.

Generally speaking, system filtration concepts are not designed to adequately deal with large quantities of dirt as occur in connection with:
- Component machining
- System assembly
- System filling
- System repair work

A study conducted by the University of Hanover describes the factors impacting the fatigue life of roller bearings as follows: "The quantity of contamination in the lubricant is described by the particle quantity and size. Combining this with particle hardness and geometry results in the type and extent of damage to raceways, with the extent also being affected by the elasto-plastic behavior of the material. The amount of damage is determined by the quantity of particles in the lubrication gap and the rollover frequency. Continued rollover leads to cracking, which in the form of fatigue damage (pitting) leads to roller bearing damage (bearing failure)."

In practice ball bearings with their punctiform contact are shown in most cases to be less sensitive to particulate contamination than roller bearings with their linear contact. Friction bearings with their larger lubrication gaps are the least sensitive to particulate contamination.
The table to the right provides an overview of the most common gap sizes illustrated in Figure 7.

Comprehensive studies of particle distributions on components and in hydraulic systems have shown that at the beginning of a system’s life, i.e. during assembly and commissioning, the particles are larger than during subsequent operation.

These large particles – up to several millimeters in size in part – can cause spontaneous outages: valve blockages, substantial preliminary damage to pumps, and destruction of seals and gaskets followed by leakage.

Active contamination management enables this rate can be reduced and costs accordingly cut, i.e.:
- Costs caused by production stops
- Costs caused by delays in commissioning systems
- Costs incurred by longer testing periods since a flushing cycle is required to remove integral contamination
- Warranty costs
- Reworking costs

Contamination management counters the situation as follows: In new systems the individual components are brought to a uniform cleanliness level, the filling fluid is kept at a defined level, as is the fluid during system operation.

<table>
<thead>
<tr>
<th>Component</th>
<th>Typical Critical Clearance (µm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Gear Pump (J1, J2)</td>
<td>0.5 - 5</td>
</tr>
<tr>
<td>2. Vane-cell Pump (J1)</td>
<td>0.5 - 5</td>
</tr>
<tr>
<td>3. Piston Pump (J2)</td>
<td>0.5 - 1</td>
</tr>
<tr>
<td>4. Control Valve (J1)</td>
<td>5 - 25</td>
</tr>
<tr>
<td>5. Servo Valve (J1)</td>
<td>5 - 8</td>
</tr>
</tbody>
</table>

Figure 7

- Destroyed raceway of a ball bearing caused by particulate contamination
- Embedded in the surface of a friction bearing
This international standard describes the gravimetric method for determining the particulate contamination of hydraulic fluids.

The objective of the procedures described below is to enable a reproducible classification of particulate contaminants in fluids. Currently there are 4 procedures for classifying particulate contaminants in fluids: ISO 4405, ISO 4406:1999, NAS 1638, SAE AS 4059 (see chart below)

<table>
<thead>
<tr>
<th>Standard</th>
<th>ISO 4405</th>
<th>ISO 4406:1999</th>
<th>NAS 1638</th>
<th>SAE AS 4059</th>
</tr>
</thead>
<tbody>
<tr>
<td>Application</td>
<td>Highly contaminated media, e.g. washing media, machining fluids</td>
<td>Hydraulic fluids</td>
<td>Hydraulic fluids</td>
<td>Hydraulic fluids</td>
</tr>
<tr>
<td>Parameters</td>
<td>Number of particles (mg/liters of fluid)</td>
<td>Number of particles</td>
<td>Number of particles</td>
<td>Number of particles</td>
</tr>
<tr>
<td></td>
<td>&gt; 4 µm(c)</td>
<td>5 - 15 µm</td>
<td>&gt; 4 µm(c)</td>
<td>&gt; 4 µm(c)</td>
</tr>
<tr>
<td></td>
<td>&gt; 6 µm(c)</td>
<td>5 - 25 µm</td>
<td>&gt; 6 µm(c)</td>
<td>&gt; 6 µm(c)</td>
</tr>
<tr>
<td></td>
<td>&gt; 14 µm(c)</td>
<td>25 - 50 µm</td>
<td>&gt; 14 µm(c)</td>
<td>&gt; 14 µm(c)</td>
</tr>
<tr>
<td></td>
<td>&gt; 21 µm(c)</td>
<td>50 - 100 µm</td>
<td>&gt; 21 µm(c)</td>
<td>&gt; 21 µm(c)</td>
</tr>
<tr>
<td></td>
<td>&gt; 38 µm(c)</td>
<td>&gt; 100 µm</td>
<td>&gt; 38 µm(c)</td>
<td>&gt; 38 µm(c)</td>
</tr>
<tr>
<td></td>
<td>&gt; 70 µm(c)</td>
<td></td>
<td>&gt; 70 µm(c)</td>
<td></td>
</tr>
</tbody>
</table>

1. **Manual evaluation:**
The fluid undergoing analysis is filtered through a prepared membrane and the cleanliness class (contamination rating) estimated or counted by hand using a microscope.

2. **Automated particle counting:**
The fluid undergoing analysis is conducted through a particle counter, which tallies the particle fractions.

This international standard describes the gravimetric method for determining the particulate contamination of hydraulic fluids.

**Basic principle**
A known volume of fluid is filtered through one or two filter disks using vacuum action and the weight differential of the filter disks (upstream and downstream of filtration) measured. The second membrane is used for evaluating accuracy.

In order to determine the gravimetric contamination of the fluid, a representative sample has to be taken from the system. ISO 4405 describes the cleaning procedure for the equipment being used. It also describes the preparatory procedures for the analysis membranes.

The membranes are flushed with isopropanol prior to use, dried in a drying oven until they achieve a constant weight, and then cooled in a defined dry environment. It is important that cooling takes place in a defined dry environment, otherwise the membrane absorbs moisture from the surroundings, thus skewing the final result.

Afterwards the membrane is weighed and this value recorded as m (T).

The membranes are then fixed in the membrane retainer and the fluid undergoing analysis is filtered. This is followed by flushing off the contaminant on the membrane using filtered solvent to completely remove the contaminant. When analyzing oil-laden fluids it is important that the remaining oil is completely flushed off the membrane.

This is followed by drying the membrane, cooling, and weighing it (as described above). The measured value is now recorded as m (E).

Gravimetric contamination is calculated as follows: \( M(G) = m(E) - m(T) \)
In ISO 4406, particle counts are determined cumulatively, i.e. > 4 µm (c), > 6 µm (c) and > 14 µm (c) (manually by filtering the fluid through an analysis membrane or automatically using particle counters) and allocated to measurement references.

The goal of allocating particle counts to references is to facilitate the assessment of fluid cleanliness ratings.
Like ISO 4406, NAS 1638 describes particle concentrations in liquids. The analysis methods can be applied in the same manner as ISO 4406:1987.

In contrast to ISO 4406, certain particle ranges are counted in NAS 1638 and attributed to measurement references.

The following table shows the cleanliness classes in relation to the particle concentration analyzed.

<table>
<thead>
<tr>
<th>Particle Size (um)</th>
<th>5-15</th>
<th>15-25</th>
<th>25-50</th>
<th>50-100</th>
<th>&gt;100</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. of Particles in 100 ml Sample</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>00</td>
<td>125</td>
<td>22</td>
<td>4</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>0</td>
<td>250</td>
<td>44</td>
<td>8</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>1</td>
<td>500</td>
<td>89</td>
<td>16</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>2</td>
<td>1,000</td>
<td>178</td>
<td>32</td>
<td>6</td>
<td>1</td>
</tr>
<tr>
<td>3</td>
<td>2,000</td>
<td>356</td>
<td>63</td>
<td>11</td>
<td>2</td>
</tr>
<tr>
<td>4</td>
<td>4,000</td>
<td>712</td>
<td>126</td>
<td>22</td>
<td>4</td>
</tr>
<tr>
<td>5</td>
<td>8,000</td>
<td>1,425</td>
<td>253</td>
<td>45</td>
<td>8</td>
</tr>
<tr>
<td>6</td>
<td>16,000</td>
<td>1,850</td>
<td>506</td>
<td>90</td>
<td>16</td>
</tr>
<tr>
<td>7</td>
<td>32,000</td>
<td>5,700</td>
<td>1,012</td>
<td>180</td>
<td>32</td>
</tr>
<tr>
<td>8</td>
<td>64,000</td>
<td>11,600</td>
<td>2,025</td>
<td>360</td>
<td>64</td>
</tr>
<tr>
<td>9</td>
<td>128,000</td>
<td>22,800</td>
<td>4,050</td>
<td>720</td>
<td>128</td>
</tr>
<tr>
<td>10</td>
<td>256,000</td>
<td>45,600</td>
<td>8,100</td>
<td>1,440</td>
<td>256</td>
</tr>
<tr>
<td>11</td>
<td>512,000</td>
<td>91,200</td>
<td>16,200</td>
<td>2,880</td>
<td>512</td>
</tr>
<tr>
<td>12</td>
<td>1,024,000</td>
<td>182,400</td>
<td>32,400</td>
<td>5,760</td>
<td>1,024</td>
</tr>
</tbody>
</table>

Increasing the class by 1 causes the particle count to double on average.

The particle counts of class 10 are bold-faced in the above table.

*Figure 11. Microscopic Examination of an Oil Sample (100 ml) Magnification 100x (NAS 10)*
Like ISO 4406 and NAS 1638, SAE AS 4059 describes particle concentrations in liquids. The analysis methods can be applied in the same manner as ISO 4406:1999 and NAS 1638.

The SAE cleanliness classes are based on particle size, number and distribution. The particle size determined depends on the measurement process and calibration; consequently the particle sizes are labeled with letters (A - F).

The SAE cleanliness classes can be represented as follows:

1. Absolute particle count larger than a defined particle size
   **Example:** **Cleanliness class according to AS 4059:**
   The maximum permissible particle count in the individual size ranges is shown in the table in boldface.
   Size B particles may not exceed the maximum number indicated for class 6.
   6 B = max. 19,500 particles of a size of 5 µm or 6 µm (c)

2. Specifying a cleanliness class for each particle size
   **Example:** **Cleanliness class according to AS 4059:**
   Size B (5 µm or 6 µm (c)): 38,900 particles / 100 ml
   Size C (15 µm or 14 µm(c)): 3,460 particles / 100 ml
   Size D (25 µm or 21 µm(c)): 306 particles / 100 ml

3. Specifying the highest cleanliness class measured
   **Example:** **Cleanliness class according to AS 4059:**
   The 6 B – F specification requires a particle count in size ranges B – F. The respective particle concentration of cleanliness class 6 may not be exceeded in any of these ranges.

<table>
<thead>
<tr>
<th>Size ISO 4402 Calibration or Visual Counting</th>
<th>&gt; 1 µm</th>
<th>&gt; 5 µm</th>
<th>&gt; 15 µm</th>
<th>&gt; 25 µm</th>
<th>&gt; 50 µm</th>
<th>&gt; 100 µm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Size ISO 11171, Calibration or Electron Microscope**</td>
<td>&gt; 4 µm(c)</td>
<td>&gt; 6 µm(c)</td>
<td>&gt; 14 µm(c)</td>
<td>&gt; 21 µm(c)</td>
<td>&gt; 38 µm(c)</td>
<td>&gt; 70 µm(c)</td>
</tr>
<tr>
<td>Size Coding A</td>
<td>B</td>
<td>C</td>
<td>D</td>
<td>E</td>
<td>F</td>
<td></td>
</tr>
<tr>
<td>000</td>
<td>195</td>
<td>76</td>
<td>14</td>
<td>3</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>00</td>
<td>390</td>
<td>152</td>
<td>27</td>
<td>5</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>0</td>
<td>780</td>
<td>304</td>
<td>54</td>
<td>10</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>1</td>
<td>1,560</td>
<td>609</td>
<td>109</td>
<td>20</td>
<td>4</td>
<td>1</td>
</tr>
<tr>
<td>2</td>
<td>3,120</td>
<td>1,220</td>
<td>217</td>
<td>39</td>
<td>7</td>
<td>1</td>
</tr>
<tr>
<td>3</td>
<td>6,250</td>
<td>2,430</td>
<td>432</td>
<td>76</td>
<td>13</td>
<td>2</td>
</tr>
<tr>
<td>4</td>
<td>12,500</td>
<td>4,860</td>
<td>864</td>
<td>152</td>
<td>26</td>
<td>4</td>
</tr>
<tr>
<td>5</td>
<td>25,000</td>
<td>9,730</td>
<td>1,730</td>
<td>306</td>
<td>53</td>
<td>8</td>
</tr>
<tr>
<td>6</td>
<td>50,000</td>
<td>19,500</td>
<td>3,460</td>
<td>612</td>
<td>106</td>
<td>16</td>
</tr>
<tr>
<td>7</td>
<td>100,000</td>
<td>38,900</td>
<td>6,920</td>
<td>1,220</td>
<td>212</td>
<td>32</td>
</tr>
<tr>
<td>8</td>
<td>200,000</td>
<td>77,900</td>
<td>13,900</td>
<td>2,450</td>
<td>424</td>
<td>64</td>
</tr>
<tr>
<td>9</td>
<td>400,000</td>
<td>156,000</td>
<td>27,700</td>
<td>4,900</td>
<td>848</td>
<td>128</td>
</tr>
<tr>
<td>10</td>
<td>800,000</td>
<td>311,000</td>
<td>55,400</td>
<td>9,800</td>
<td>1,700</td>
<td>256</td>
</tr>
<tr>
<td>11</td>
<td>1,600,000</td>
<td>623,000</td>
<td>111,000</td>
<td>19,600</td>
<td>3,390</td>
<td>1,020</td>
</tr>
<tr>
<td>12</td>
<td>3,200,000</td>
<td>1,250,000</td>
<td>222,000</td>
<td>39,200</td>
<td>6,780</td>
<td></td>
</tr>
</tbody>
</table>

Table shows the cleanliness classes in relation to the particle concentration determined.

*Particle sizes measured according to the longest dimension.

**Particle sizes determined according to the diameter of the projected area-equivalent circle.
A representative sample is taken of the fluid and analyzed as follows:

1. **Manual procedure according to ISO 4407** (Hydraulic fluid power – Fluid contamination – Determination of particulate contamination by the counting method using a microscope).

ISO 4407 contains a description of a microscopic counting method for membranes. 100 ml of the sample undergoing analysis is filtered through an analysis membrane featuring an average pore size of < 1 µm and square markings.

The standard also describes the cleaning procedure and maximum particle count of the negative control. After the analysis membranes are dried, 10, 20 or 50 squares are counted depending on the size of the particles, followed by adding the values and extrapolating to the membrane diameter.

The manual count of the particles is done in the “old” levels of > 5 µm and > 15 µm since the longest dimension of a particle is counted in ISO 4407 yet the diameter of the area-equivalent circle is counted in the “new” ISO 4406:1999. As described above, the reference values obtained for this count correspond to the reference values of the “new” evaluation.

This counting method can only be used for very clean samples. Generally speaking, the cleanliness classes are estimated on the basis of reference photographs or the samples automatically counted.

2. **Automated particle counting**

Below follows a description of how common particle counters employing the light extinction principle function.

The figure left shows a simplified rendering of the measurement principle employed in the light extinction principle.

The light source transmits the light (monochromatic light for the most part) onto an optical sensor, which emits a specific electrical signal. A shadow is created on the photodiode if a particle (black) comes between the light source and the photodetector. This shadow causes the electric signal emitted by the sensor to change. This change can be used to determine the size of the shadow cast by this particle and thus the particle size to be determined.

This procedure enables the cleanliness classes according to ISO 4406:1987, ISO 4406:1999, NAS 1638 and SAE AS 4059 to be accurately determined.

The “noise” involved in this measurement principle is extraneous liquids and gases which cause the light beam to be interrupted and thus be counted as particles. The particle counter should be calibrated according to ISO 11943 (for ISO 4406:1999).
Determining the residual dirt quantities present on components can be done employing quantitative and qualitative factors.

**Quantitative:**
- mg/component
- mg/surface unit (oil-wetted surface)
- mg/kg component weight
- no. of particles > x µm/component
- no. of particles > x µm/surface unit (oil-wetted surface)

**Qualitative:**
- length of the largest particle (subdivision into hard/soft)

Components with easily accessible surfaces are components in which only the outer surface is of interest for the most part when performing residual dirt analyses. There are exceptions here, e.g. transmission and pump housings, as the internal surface of interest. These components belong to group 1 and their surfaces are not easily accessible in most cases.

Components in which the inner surfaces are examined or preassembled assemblies belong to group 2; for the analysis procedure for this group, refer to page 16.

There are two methods that can be used to determine the residual dirt of group 1 components.

**Ultrasonic Method**

The ultrasound method involves submitting the components to an ultrasonic bath, exposing them for a defined period of time at a defined ultrasonic setting and bath temperature. The particulate contamination is loosened by the exposure and then flushed off the component using a suitable liquid.

The particle dispersion in the flushing liquid obtained in this manner is analyzed according to specified evaluation methods.

The ultrasonic energy setting and the duration of exposure have to be indicated in reporting the result. The ultrasonic procedure is particularly suitable for small components in which all surfaces have to be examined. **Cast components and elastomers should not be subjected to ultrasonic washing if possible a risk is posed here of the carbon inclusions in the cast piece being dissolved, thus skewing the results. These effects have to be ascertained prior to performing an ultrasonic analysis.**

Components with easily accessible surfaces or components in which only surface parts have to be examined are analyzed using the flushing method. This method involves flushing the surface undergoing analysis in a defined clean environment using an analysis fluid, which also has a defined cleanliness. A “negative control” or basic contamination control is performed prior to analysis in which all the surfaces of the environment, e.g. the collecting basin, are flushed and the value obtained reported as the basic contamination of the analysis equipment. The flushing fluid is then analyzed using the specified evaluation methods.

The darker areas to the right are the flushing areas; those to the left and lighter are the designated analysis area. In reality these two circuits are configured using suitable valves in such a manner that switchover can be done between the two storage tanks. The figure represents a simplified circuit diagram. The analysis fluid is subjected to a pressure of approximately 58 – 87 psi (4 – 6 bar) and conveyed through the system filter and the spray gun into the analysis chamber. The system filter ensures that the analysis fluid sprayed on the surface being examined has a defined cleanliness. The particle-loaded fluid collects in the collecting basin and is filtered through the analysis membrane via vacuum action. The membrane is then evaluated according to the analysis methods described on the following pages.

This method is very rarely used, as it is very difficult to reproduce manually. However, results are reproducible when automatic shakers such as those used in chemical laboratories are employed. The analyzed components are components subject to wear whose inner surfaces are to be analyzed (e.g. pipes, tanks). The important thing is that the particles are flushed out of the inside of the components after being shaken.

The table on the following page shows a comparison of the various methods for analyzing components and assemblies.
### Flushing Method

**How Performed**
Components are flushed with the analysis fluid in a defined clean environment.

**Applications**
Components in which only surface parts have to be examined and components in which ultrasound may damage the surfaces. Components with a simple design and with easily accessible surfaces.

**Pros**
Analysis can be performed quickly

**Cons**
Standards are not yet available (currently in preparation)

### Ultrasonic Method

**How Performed**
Components are exposed to an ultrasonic bath and are then flushed with the analysis fluid.

**Applications**
Small components and components in which all surfaces are to are to be analyzed (the component size depends on the ultrasonic bath).

**Pros**
Reproducibility

**Cons**
Analysis takes a long time

---

### Evaluation Methods

Evaluating particle-laden flushing fluids can be done according to various criteria. Gravimetric analysis is useful for heavily contaminated components, whereas particle counts in various size ranges are useful for very clean components.

The following table provides an overview of the individual evaluation methods.

<table>
<thead>
<tr>
<th>Manual Methods</th>
<th>Automated Methods</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Gravimetric method</strong> [mg/m²]</td>
<td>Counting of particles on the analysis membrane [no. of particles &gt; x µm/m²]*</td>
</tr>
<tr>
<td>The particle-laden fluid is filtered through a prepared analysis membrane</td>
<td>Counting of particles on the analysis membrane [no. of particles &gt; x µm/m²]*</td>
</tr>
<tr>
<td>The analysis membrane is weighed before and after analysis and the gravimetry is computed on the basis of the difference between the measured values</td>
<td>The analysis membrane is placed under a microscope and evaluated using a software tool. This software records the light-dark contrasts on the membrane and interrupts them as particles.</td>
</tr>
<tr>
<td>Samples exhibiting contamination &gt;10 mg</td>
<td>Samples featuring high a content of coarse contamination. Often combined with gravimetric evaluation.</td>
</tr>
<tr>
<td>ISO 4405</td>
<td>ISO 4407</td>
</tr>
</tbody>
</table>

**Advantages**
- Can be used for large particle quantities
- Measurement range selectable (2-400µm).
- Accurate measurement method

**Disadvantages**
- Takes a long time (min. 1 h)
- Lab Method

**Applications**
- On-line process control in manufacturing and assembly.
- Can also be used in labs.
The following table provides an overview of applications of the analysis and evaluation methods.

<table>
<thead>
<tr>
<th>Evaluation Method</th>
<th>Gravimetry</th>
<th>Particle Counting</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Analysis Method</td>
<td>Flushing</td>
</tr>
<tr>
<td>Simple Components</td>
<td>easy-to-access surfaces; gears</td>
<td>U</td>
</tr>
<tr>
<td>Components</td>
<td>internal surfaces pipes, tanks</td>
<td>U</td>
</tr>
<tr>
<td>Complex Components</td>
<td>components featuring various bore holes or ducts; control plates</td>
<td>CU*</td>
</tr>
<tr>
<td>Simple Systems</td>
<td>surface is to be analyzed immersed sensors</td>
<td>U</td>
</tr>
<tr>
<td>Systems</td>
<td>internal surfaces rails of common rail systems</td>
<td>CU*</td>
</tr>
<tr>
<td>Complex Systems</td>
<td>valves, pumps</td>
<td>CU*</td>
</tr>
</tbody>
</table>

* It has to be ensured that the particles dislodged from the component can be flushed away.

U = Usable
CU = Conditionally usable
NU = Not usable

SMART® (Schroeder’s Micronic Automated Recirculating Technology) utilizes a microprocessor to customize operations based on user input. Products that use SMART® technology are marked throughout the catalog with the SMART® Product logo.

Schroeder’s EPK Patch Test Kit (shown to the right) provides the tools needed to pull contaminated fluid through a patch and compare the resulting patch under a microscope to representative photos of various contamination levels to determine the fluid’s ISO level.
The cleanliness of components and systems that pass through a flushing or test stand can be determined on the basis of the cleanliness of the test fluid in some cases. This indirect analysis method is preceded by manual analyses for validation purposes.

For example, hoses are flushed by hand and the results evaluated in accordance with the methods discussed on page 13. At the same time, the cleanliness of the test fluid of the test stand is determined in the return flow, i.e. downstream of the component.

If a correlation is detected between the manual and the automatic (indirect) value, this means that indirect value analysis can be selected as a measure of quality.

The flushing stand used for analyzing the residual dirt content of systems has to feature the following:
1. Flushing has to be done using as turbulent a flow as possible.
2. The fluid used has to possess a dispersion effect.
3. All channels and surfaces have to be exposed to the flow.
4. The effectiveness of flushing can be improved by pulsating the flushing.

The Reynolds number – a dimensionless reference – describes the flow properties of fluids. Below follows a brief description of how the Reynolds number is derived using pipe flow as an example.

Weights are discounted in the calculation of the Reynolds number. Generally speaking, only pressure, friction and inertial forces affect fluid elements and bodies subjected to flows. They have to be in balance at all points of the flow. If the relationship of friction and inertial forces is the same in similar points P1 and P2, then similar flows are said to be present.

The Reynolds equation looks like this when the above properties are taken into account:

\[ Re = \frac{\text{mean velocity} \times \text{internal pipe diameter}}{\text{kinematic viscosity}} \]

\[ Re = 21220 \times \frac{Q}{d_i \times V} \]

Whereby: \( Q \) = volumetric flow rate (l/min), \( V \) = viscosity (mm²/s), and \( d_i \) = inside pipe diameter (mm).

The critical Reynolds number \( Re_{\text{crit}} \) depends on kinematic velocity \( V \), flow rate \( Q \) of the fluid, and the geometry of the passage through the flow is being conducted. If the Reynolds number of a flow is smaller than \( Re_{\text{crit}} \), the flow is said to be laminar. Turbulent flow is said to be present for values above \( Re_{\text{crit}} \). The critical Reynolds number for oil is given below.

\[ Re_{\text{crit}} \text{ oil} = 1900 \text{ – } 3000 \]

(Source: Kahrs, M.: Der Druckverlust in Rohrleitungen Ölhydraulischer Antriebe; VDI Forschungsheft 537, Düsseldorf 1970)

The diagram to the left shows a parabolic, laminar flow in a pipe. This shows that the flow velocity of a laminar current in the middle of the pipe (peak of the parabola) is larger than along the pipe wall.

In a turbulent flow this parabola flattens and spreads (when mean values are considered), as transversal currents are involved in a turbulent flow. They cause the flow velocity to be increased in the vicinity of the pipe walls.

This effect is utilized when flushing systems as increasing the flow rate causes particles that have been deposited on the wall to be loosened and swept away.

**Analysis of the Cleanliness of Systems on a Flushing/Test Stand**

**Turbulent Flow**

**Difference Between Laminar and Turbulent Flow**

**Figure 16. Similar Flows Around Different Cylinders**

**Figure 17. Laminar Flow**
All particles move without mixing.

The path of a particle is described by a stream thread.

Parabola-shaped velocity distribution (applies to pipes).

Reynolds number smaller than \( Re_{\text{crit}} \)

**Figure 18. Turbulent Flow**
All particles are continuously mixed.

The path of a particle cannot be predicted.

Relatively even time-averaged velocity distribution (flattened parabola).

Reynolds number larger than \( Re_{\text{crit}} \)

Source: University of Würzburg Fluid Mechanics lecture
The oil used for flushing has to have a dispersion effect so that particles are dislodged and transported off. Special thin-bodied mineral-oil-type flushing oils can contribute instrumentally to improving the flushing effect. They lower the adhesion force between the dirt particles and the pipe wall. By virtue of their excellent surface wetting properties, they creep into between the dirt particles and the wall, thus causing the particles to become dislodged. Experiments have shown that by changing the flushing fluid from an operating fluid to a flushing oil, component/system cleanliness can be increased by a factor of 4. Flushing oils of this type have to then be closely matched to the hydraulic medium used as failure to properly match the two may lead to marked foaming, filter blockage and clogging of the system.

When setting up the inspection and testing plan, it has to be ensured that all surfaces and ducts are wetted during flushing.

The cleanliness of components and systems which undergo function testing can be determined on a flushing or function test stand (= flushing stand).

This method is used for pumps, cylinders, transmissions, control units, power steering units, valve blocks, etc.

Once it is ensured that the flushing stand possesses the properties indicated above, an analysis is conducted as described below.

Prior to the analysis the flushing stand is cleaned to a defined high cleanliness level so that the basic contamination of the test system does not affect the measurement results. Then this basic cleanliness is computed and recorded.

The sampling site for an automatic particle counter is defined as a site upstream or downstream of the test item, which is subjected to a direct flow. The following is performed if the analysis result is to be additionally subjected to gravimetric analysis: the entire test fluid is collected and filtered through an analysis membrane or an inline membrane retainer featuring the analysis membrane is integrated in the return-flow line.

Now the test item is tested in accordance with the inspection and testing plan, during which the cleanliness classes are recorded.

Example 1: The schematic below shows the analysis performed on a pump test stand.

Example: As-supplied condition:
17 / 15 / 12 according to ISO 4406:1999

1. Warning point: 18 / 16 / 13 for 3 successive measurements
2. Stop signal: When exceeding 18 / 16 / 13 limit cleanliness class in 2 successive measurements.

After 5 minutes of testing the pump speed, it is increased to the maximum speed. This causes particulate contamination to be dislodged. The system becomes increasingly cleaner. Particulate contamination is still being released after 1 hour of testing (standard test time: 10-15 min.), consequently the cleanliness of the return-line fluid (blue = downstream of the test item) never achieves the same cleanliness as upstream of the test item.

This method is suitable for checking the cleanliness of items being delivered quickly and simply in series testing, documenting it and then concluding the flushing procedure when the target value is achieved. By integrating the measurement circuit in the manufacturing instrumentation and control system it is also possible to quickly detect any deviations and initiate suitable measures. The goal of continuous cleanliness monitoring is to monitor process reliability with regard to system cleanliness upon delivery.

A specification like this also enables increased system contamination to be responded to quickly. If these measurements are only conducted once a day, a whole day’s output might be affected and have to be remedied. The result is unnecessary costs that can be avoided by integrating a continuous measurement procedure.

When conducting a reference measurement, the system is disassembled after the test run, if possible, and the individual components analyzed using the flushing method.
The reliability of hydraulic systems can be impacted heavily by particulate contamination during the running-in phase. The risk of outages during the first minutes or hours of operation is particularly high as the foreign particles introduced or created during the assembly process are still relatively large and can thus cause sudden outages. During continued operation, these large particles are ground into smaller ones, the result being that damage can be caused to the surfaces of system components during this crushing process. The consequences are leakage, degraded output and efficiency, or a shortening of the component's service life. In many cases, microfiltering is used to quickly clean the system fluid during commissioning. This is where contamination monitoring is key in the manufacture and assembly of these systems. By implementing contamination management a major portion of particulate contamination introduced during manufacture and assembly can be removed. The result is cost savings by virtue of smaller performance deviations on test stands caused by the sudden clogging of particles in sensitive system components plus lower costs associated with warranty and non-warranty courtesy work. For more information, refer to page 28.

Below follows a description of the goal, design and performance of a process audit.

Contamination monitoring extends to checking the cleanliness status of all manufacturing and assembly processes considered relevant in this connection. Proper preparation and informing all those involved are key in contamination monitoring.

First, the objective of contamination monitoring is specified, e.g.
- Determining the current situation
- Checking fluctuations between batches
- Checking washing processes
- Comparing the target with the actual situation
- Determining the sampling point

During the planning and design phase, the sampling points for components and taking liquid samples are determined using a production plan or operation sheet. The employees to be involved in contamination monitoring are informed of the objectives and procedures.

NOTE:
Manufacturing has to continue in the same manner, meaning that no additional cleanliness levels, etc. are to be integrated. The purpose of contamination monitoring is not to check the quality produced by the employees but rather determining the causes and sources of contamination.

The schematic above shows the manufacturing processes and the corresponding sampling points. However, in actuality sampling is more comprehensive, i.e. the description includes the number of the Minimess fittings at which sampling is done, for example.

A representative sampling is taken of the fluids and components; the samples are stored so as to prevent any further contamination. Special sampling bottles are used for the fluid samples; the components are stored in defined clean packaging.

The analysis is performed in accordance with the methods specified on pages 13 - 17 and the findings recorded.

Properly trained or experienced individuals while inspecting the manufacturing and assembly line can detect some sources of contamination. That is why such an inspection is conducted during the audit. The findings made during inspection are then compared with the results in hand.
The contamination monitoring results describe the condition at the time at when sampling is done. The findings might look like this:

**Figure 21. Housing Processing**

This chart shows an excerpt of the housing manufacturing process. The component samples are taken upstream and downstream of the washing station. The findings show that the washing station performs well and that it is well positioned here. Subsequent storage is not being done properly as the portion of particulate contamination is almost double.

By applying a cleanliness specification to components and the system it can be ensured that as-supplied quality is constant.

The following should be in mind when drafting a cleanliness specification:

- State of the art
- Benchmarking – what do others do?
- Inclusion of previous experience – if available
- Defining and implementing contamination management as an “official project”
- Inclusion of all hierarchy levels
- Accurate documentation of how the specification was developed
- Developing clear-cut definitions

Next, it has to be determined which components in the system are the most sensitive. Frequently it is not possible to achieve the same level of cleanliness throughout the system during assembly. If suitable filtration takes place upstream of the sensitive components, an area of low-contamination-sensitive components can be defined upstream of this filtration and an area of highly contamination-sensitive components downstream of the filter.

These individual components or system areas should be subdivided into sensitivity areas.

<table>
<thead>
<tr>
<th>Category</th>
<th>Designation</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Low particle-sensitivity</td>
<td>For the most part low-pressure systems with large gap tolerances</td>
</tr>
<tr>
<td>B</td>
<td>Particle-sensitive</td>
<td>Low-pressure systems with small gap tolerances</td>
</tr>
<tr>
<td>C</td>
<td>High particle sensitivity</td>
<td>High-pressure systems with small gap tolerances and with exacting demands made of safety and security systems</td>
</tr>
</tbody>
</table>

A maximum particulate contamination value is specified for each of these cleanliness categories.

A car motor illustrates this subdivision below:

<table>
<thead>
<tr>
<th>Category</th>
<th>Motor Area</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Air /Coolant water circuit</td>
</tr>
<tr>
<td>B</td>
<td>Low-pressure oil circuit</td>
</tr>
<tr>
<td>C</td>
<td>Diesel direct injection / High-pressure oil circuit</td>
</tr>
</tbody>
</table>

In addition, the fluid cleanliness ratings of the individual system and process fluids are defined.
The following parameters are defined in the cleanliness specifications for the components:

1. Goal of the cleanliness specification
2. Applicability (system designation)
3. Extent of inspection and testing; inspection and testing cycles
4. Sampling
5. Analysis method
6. Evaluation method
7. Accuracy
8. Analysis fluids to be used
9. Documentation
10. Limit values

This specification has to be made for each individual system; consequently a few things are discussed which have to be borne in mind.

Work instructions concerning sampling, analysis and evaluation methods should be described in detail so as to ensure that sampling is always done in a uniform manner. In addition, the analysis results depend on the analysis fluid and method, particularly when it comes to component analysis. Documentation should be done using forms so that all the results are readily accessible.

### Example of a Cleanliness Specification

1. **Goal of the cleanliness specification**
   The goal in implementing this cleanliness specification is to achieve a constant level of cleanliness for system X.

2. **Applicability (system designation)**
   This specification applies to system X including its series A, B, and C. It extends to all components whether sourced or manufactured in house. It also specifies the system fluids of system X with regard to their cleanliness.

3. **Extent of inspection & testing; inspection & testing cycles**
   5 samples a month of each component are to be taken and analyzed. If the supplier parts achieve a constant cleanliness value after 6 months, the sampling cycle can be extended to every 2 or 3 months. An analysis of the entire (assembled) system is to be done at least once a week prior to delivery. Checking of the fluid cleanliness should be done on a continuous basis.

4. **Sampling**
   Sampling of components is to be done at goods receiving and is to be representative. Samples should be packed in a dust-tight manner and sent into the laboratory. The fluid samples are to be taken at the sampling points indicated in the inspection and testing plan.
5. Analysis method
The flushing method should be used for component analysis. The surfaces of the component are flushed in a clean environment using x ml of the test fluid (XY) which has a cleanliness of xx, under a pressure of z bars as specified by the inspection and testing plan. The flushed-off particulate contamination is collected on an analysis membrane and subjected to gravimetric analysis. Representative samples are taken of the system fluids at the specified sampling points. All testing parameters are specified; the duration of testing, what is tested, the pressures, and speeds. When conducting static inspection and testing make sure that a flushing effect is present so that the cleanliness of these components can be determined, (the static pressure test has to be followed by a dynamic flushing process in order to analyze the actual quantity of particles which is flushed out of the component.)

6. Evaluation method
In the component analyses the analysis membrane is dried until it achieves a constant weight, and then cooled in a defined dry environment and weighed. This procedure is repeated subsequent to filtration. The weight differential indicates the “gravimetric contamination” of the component. This is followed by visually examining the analysis membranes through a microscope and measuring the longest particles. Evaluation of the fluid samples is done in accordance with ISO 4405, ISO 4407, ISO 4406:1999 or NAS 1638.

7. Accuracy
The analysis equipment has to be brought to a residual dirt content of 0.2 mg prior to conducting the analysis so that the measurements taken of the component samples are sufficiently accurate. This is determined by performing a negative control, i.e. flushing the equipment without testing. When the result of the analysis drops below 0.5 mg, the batch size is to be increased and thus a mean value of the results computed.

8. Analysis fluids to be used
The following analysis fluid should be used for the component analyses: ABC-XX, with a cleanliness class of 14 / 12 / 9 and no particles > 40 µm.

9. Documentation
The documentation of the results is done using a result sheet.

10. Limit values
The components are subdivided into 3 cleanliness classes:

<table>
<thead>
<tr>
<th>Category</th>
<th>Designation</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
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<tr>
<td>C</td>
<td>High particle sensitivity</td>
<td>High-pressure systems with small gap tolerances and exacting demands</td>
</tr>
</tbody>
</table>

The following cleanliness specifications apply to each of these classes (fictitious example).

<table>
<thead>
<tr>
<th>Category</th>
<th>Gravimetry</th>
<th>Particle Sizes</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>20 mg / component</td>
<td>Max. 4 particles &gt; 500 µm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Max. size: 400 µm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>No fiber bundles</td>
</tr>
<tr>
<td>B</td>
<td>10 mg / component</td>
<td>Max. 4 particles &gt; 400 µm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Max. size: 800 µm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Fibers up to 4 mm</td>
</tr>
<tr>
<td>C</td>
<td>5 mg / component</td>
<td>Max. 4 particles &gt; 200 µm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Max. size: 1,000 µm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Fibers up to 2 mm</td>
</tr>
</tbody>
</table>

The transmission components are subdivided into the individual categories.

Group A: crankcase sump.
Group B: intermediate housing, transmission housing, coupling flange
Group C: valve plate, valve housing, centering plate

Fluid samples:
At the end of the test run, the transmission fluid may not fall short a cleanliness rating of 17 / 15 / 13 (c) according to ISO 4406:1999. The system is to be operated using a cleanliness rating of 18 / 16 / 14 (c) according to ISO 4406:1999.

11. Procedure to be followed in the event that the specification is not adhered to
The supplier components are to be returned to the supplier in the event that the specification is not adhered to. If this procedure results in production delays, the components will be cleaned and analyzed by us at the supplier’s expense.
Sources of Contamination in the Manufacturing and Assembly of Hydraulic Systems

Particulate contamination can enter a fluid power system in various ways. The main sources of ingestion are shown in the following diagram.

Figure 24. Sources of Contamination in the Manufacturing of Hydraulic Systems

Some of these sources of contamination can be eliminated in a simple, cost-effective manner.

The ingestion of contamination in the manufacturing and assembly of hydraulic systems can be eliminated in a cost-effective manner in various process steps.

Storage and Logistics
When storing and transporting the components and systems care has to be exercised to make sure that they are properly sealed shut or well packed. Transportation and storage packing has to be in keeping with the cleanliness status of the individual components.

Assembly of Systems and Subassemblies
The assembly of these systems is to be done in accordance with system requirements. This means that the assembly and mechanical fabrication areas have to be separated if necessary in order to prevent the ingress of contamination. The assembly stations have to be kept clean to a defined cleanliness and those working in these areas have to wear special, lint-free clothing. The assembly equipment has to be properly cleaned so as to prevent the ingress of dirt here, too.

Raising the Awareness of Employees
In order to achieve the objective of “defined cleanliness of components and systems” it is important that employees at all levels be involved in this process. Frequently, a considerable savings potential is contained in the employees’ wealth of ideas and experience — particularly those working at assembly lines and in fabrication.

Experience has shown that when employees are able to identify with the objective being striven for, they are more able to help in implementing it quickly and effectively.

Environment — Air Cleanliness
In some cases it will be necessary to set up a clean room for the final assembly of very contamination-sensitive systems, e.g. fuel systems, brakes shock absorbers, etc. This has to be decided on a case-by-case basis. However, in many cases performing the measures described here suffices.

Removal of Particulate Contamination from Hydraulic Systems (Practical Experience) and Components
Generally speaking, particulate contamination is removed from a hydraulic system via filtration. Various types of filters are used depending on the amount and type of contamination.

Belt filter systems or bag filters are used when large quantities of contaminants are involved. These filters have the job of removing the major portion of contaminants from the system. These filter types are also used for prefiltration purposes.

In most cases, these coarse filters do their job of “removing a lot of dirt from the system” very well. However, microfiltration also has to be done if a constant defined high level of cleanliness of the system fluid is to be ensured.

Whereas microfiltration ensures quality, the job of coarse filtration is to control the quantity of contamination.
Individual components are freed of clinging contamination in cleaning systems (particles, remainder of machining or corrosion protection fluids, etc.). Cleaning can be done by employing various mechanical methods (e.g. spraying, flooding, ultrasonic methods) using various cleaning fluids (aqueous solutions or organic solvents). The temperature and duration of cleaning also have a decisive effect on the cleaning effect. These factors have to be carefully matched and optimally tuned in order for a favorable cleaning effect to be achieved in an economical amount of time.

![Figure 25](image)

Various studies of washing processes have shown that some of these for the most part cost-intensive processes aren’t worthy of the name. Some people refer to washing processes as “particle distribution processes”. This “property” was detected in examinations of components sampled upstream and downstream of a washing process.

There are two possible responses in a case like this:

1. Discontinue the washing process when component cleanliness becomes worse after washing than before.
   
   **Advantage:** temporary cost savings

   **The best alternative:**

2. Optimize the process. The following should particularly be borne in mind when optimizing washing processes: cleanliness of the washing, flushing and corrosion protection fluid, mechanical aspects, suitability of the washing process for the components undergoing washing and filtration of the washing and flushing fluid.

   When purchasing washing systems, make sure to specify the component cleanliness to be achieved and the maximum contamination load of the washing fluid in terms of mg/l or a cleanliness class.

   Washing systems used to be subdivided into micro and micronic washing. This was a very imprecise definition of the cleaning performance to be achieved. Nowadays the permissible residual dirt quantity of the cleaned components is defined.

   Specifying these residual dirt quantities is done as follows: mg/component, mg/kg component, mg/surface units or particle concentrations in various size ranges. In addition, the maximum sizes of the particles are defined which can be on the washed component, e.g. max. 3 particles > 200 µm, no particles > 400 µm.

   These values cannot be achieved unless the factors indicated above are matched and fine-tuned. The following factors additionally have to be borne in mind: environmental protection and labor safety, local situation relating to space and power available, and the target throughout rate.

   The cleanliness of the washing and flushing fluids also has a decisive impact on the cleaning performance of the washing machine.

   However, we are concerned here only with the maintenance of the washing and flushing fluids.
Using Filtration as Fluid Maintenance for Separating out Particulate Contamination

Bag and backflush filters in various microfilter ratings are the standard equipment used in the maintenance of the fluid of washing systems. Although these filters are suitable for removing large quantities of contamination from a system, they are not suitable in most cases for maintaining defined cleanliness classes. Owing to their design, they do not offer much resistance, (the counterpressure built up across the filter is very low), below 15 psi for the most part. That is why this filter type is frequently used in the full flow when feeding cleaning fluid into the washing or flushing chamber. The filter housings are equipped with pressure gauges for monitoring the proper functioning of the filter.

Bag filters pose the risk that overloading can cause the bag to be destroyed and large contaminant quantities released. That is why it is advisable to additionally define minimum change intervals and to regularly monitor the cleanliness of the washing fluid in addition to the standard parameters like pH value or microbial count.

Residual dirt values of cleaned components are increasingly being defined and specified as an acceptance criterion for the cleaning system. It is of paramount importance that constant adherence be maintained to these values. It is also imperative that the quality of the cleaning fluid be maintained at a high, constant level.

This can be achieved by use of the targeted microfilters, featuring a constant and absolute separation rate. For the most part, tube filters or disk filters are used. The advantage of these filter types as compared to standard hydraulic filter elements is their high contaminant retention rate owing to their depth effect.

The high contaminant separation rate offered by these filter types removes a high amount of contamination from the washing fluid. This causes the filters to become quickly exhausted and blocked. A sufficiently long service life coupled with high washing fluid cleanliness can be achieved by combining filters for removing the main portion of contaminants from the system with absolute microfilters.

Example: At a leading automotive supplier, the camshafts were to be cleaned to a defined cleanliness of 9 mg / component. Point of departure:

Technical Specifications of the Washing Machine Present on Site

| Tank Volume: | 21 gal. (80 l) |
| Pump Delivery Rate: | 66 gpm (250 l/min) (centrifugal pump) |
| Washing Agent: | Ardox 6478 – chemetall |
| Concentration: | 2.3 – 3% |
| Bath Temperature: | ca. 122°F (50°C) |
| Filtration: | Backflush filter downstream of pump, 50 µm filter rating |

Process Data

| Bath Change Frequency: | 1 time/week |
| Throughput: | 3,000 - 4,000 |
| Wash Cycle: | 15 s/component |

Challenge: Clogging of the tank, Quality no longer sufficient after 2-3 days. Fluctuation in the contamination content of the components upstream of the line: 30 – 50 mg. Cleaning costs could not be allowed to increase, although quality still had to be improved.
Achieve a residual contaminant value of a maximum of 9 mg/camshaft
Cleanliness of washing fluid of < 30 mg/liter
Extend the service life of washing fluid, i.e. save costs associated with changing the fluid
Prevent clogging of the tank, e.g. save cleaning time
For process reliability reasons, a low-maintenance cleaning system was to result which enabled the camshafts to be cleaned to a residual contaminant content of 9 mg/component, this to be done cost-effectively.

The service life of the cleaning fluid was extended from 1 week to 8 weeks. There was no more clogging of the tank. Changing the bath fluid was done on account of the increased chloride content, not on account of contamination.

The residual contaminant values of max. 9 mg/camshaft and max. 30 mg/liter of bath fluid (when using a 5-µm membrane for analysis) were achieved and maintained at this level.

By optimizing the fluid maintenance of this washing line, an improvement in quality was achieved at no added cost and without comprising process reliability.

This example shows that prior to any such optimization or in new facilities the cleanliness of the components upstream of the system, throughput, technical details, targets have to be known and defined, for only in this way can the success of such an endeavor be ensured.

<table>
<thead>
<tr>
<th>Investment ($)</th>
<th>Recurring Costs ($)</th>
<th>Savings/Year ($)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Off-line Filtration</td>
<td>5,000.00</td>
<td></td>
</tr>
<tr>
<td>Filtration Costs</td>
<td>7,500.00</td>
<td></td>
</tr>
<tr>
<td>Extension of the Service Life of the Bath</td>
<td></td>
<td>10,000.00</td>
</tr>
<tr>
<td>Lower Reworking Costs</td>
<td>These costs can’t be quoted.</td>
<td></td>
</tr>
<tr>
<td>Down Time of the Washing Machine for Cleaning</td>
<td>These costs can’t be quoted.</td>
<td></td>
</tr>
</tbody>
</table>
Most systems come into contact with the hydraulic fluid during initial system filling or function testing. This process affords the manufacturer a substantial opportunity to impact the final cleanliness of the entire system. By using suitable filtration of the filling and test fluids, system cleanliness can be quickly optimized upon delivery or commissioning.

The cleanliness of the final product can be controlled via function testing in the same way as by a washing machine. Some companies have the following motto: “The test stand is our last washing machine.”

This statement might be true, however it is an expensive approach in practice. Yet when performing process reliability measures for supplying systems with a defined cleanliness, this is the first approach.

On a function test stand not only function testing is performed but the components and systems are run in as well. A frequent side effect of this is the flushing effect of the system undergoing testing. By employing targeted fluid maintenance and cleanliness monitoring, this flushing effect can be used to ensure that systems possess a defined, constant cleanliness status upon delivery.

Cleanliness monitoring provides information on the process stability of the upstream fabrication and cleaning steps. Frequently, continuous monitoring of test fluid cleanliness results in the cleanliness of the entire system as supplied being documented. This approach is used in mobile hydraulics, turbines or paper machinery upon delivery or during commissioning in order to demonstrate to the final customer that his system is being supplied with the specified cleanliness.

**Example:** The following study illustrates the cleaning process of a pump during commissioning:

The cleanliness of the test fluid upstream of the test item is maintained at a cleanliness rating of 16 / 14 / 11 (c). After 5 minutes of testing the pump speed is briefly increased to the maximum speed. The test run is concluded after 10 minutes.

In this case, the dirt content of the test item amounted to $1 \text{ mg/kg}$ component weight upon the conclusion of the test run.

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**Figure 29. Cleaning Process on a Pump Test Stand**

As the schematic above shows, the particle concentration continuously drops during the first 4 minutes of the test run. The particle concentration jumps when the pumps are turned up to full speed after 5 minutes. The next 5 minutes are again used for cleaning the system. Now the following can be asked: “How clean are the valves that leave this test stand?”

The flushing procedure can be monitored by occasionally disassembling the valves in a defined clean environment and evaluating the dirt content of the individual components.
Unfortunately, improper component storage is not uncommon. Seals and gaskets which arrive at the assembly line clean and packed in bags are unpacked and filled into containers which are dirty for the most part as this involves less work and effort.

In most cases, these factors are not taken into consideration and substantial savings potential that could be easily utilized through improved packaging and storage is overlooked.

Suitable cleanliness specifications for internally produced and sourced parts enable the ingress of contamination into systems to be minimized right from the beginning.

This method is most frequently chosen for large systems in order to minimize wear during commissioning.

The filtration of the flushing stand has to be designed so that during subsequent analysis the contaminants flushed out of the system undergoing testing are removed and other measurements aren’t skewed. As an alternative, cleanliness can be measured and recorded upstream and downstream of the test item during the entire measurement sequence.

In the example above, the specified sampling point was located directly downstream of the pump and an online particle counter connected.

The crane jib was extended after 6, 8 and 10 minutes. The graph clearly shows that every time a new area was brought on line contaminant sediments were flushed out.

When a system’s characteristic curve/behavior is known, cleanliness testing can be performed at the end of function testing and, thus, system cleanliness described subsequent to commissioning. This method enables process control to be implemented quickly and reliably during series testing/commissioning. The cleaning curve plotted over time is an indication of the ingress of contaminants during assembly.
The core aspects of contamination management are a cost analysis and efficiency review. The following costs are considered in the cost analysis:

- Warranty and non-warranty courtesy work
- Energy costs (e.g. cooling and reheating of washing machines during fluid changes)
- Test stand costs (test item time)
- Costs of the tools and dies of machine tools (increased wear due to high particle concentrations)
- Fluid costs (washing machines, test stations, machine tools)
- Labor costs (reworking, cleaning of washing machines, machine tools, etc.)
- Filter costs

<table>
<thead>
<tr>
<th></th>
<th>One-off Investment</th>
<th>Recurring Costs / Year</th>
</tr>
</thead>
<tbody>
<tr>
<td>Function Test Stands (5)</td>
<td>6,500 x 5 = 32,500</td>
<td>7,500 x 5 = 35,000</td>
</tr>
<tr>
<td>Storage Conditions</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Coverings for the Pallets</td>
<td>2,500</td>
<td></td>
</tr>
<tr>
<td>Washing Machine for Cleaning the Pallets</td>
<td>50,000</td>
<td>25,000</td>
</tr>
<tr>
<td>Machining Process</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Manpower/Cleaning Filtration</td>
<td>750 x 7 = 5,250</td>
<td>2,000 x 7 = 14,000</td>
</tr>
<tr>
<td></td>
<td>1,250 x 7 = 8,750</td>
<td></td>
</tr>
<tr>
<td>Consulting Expenses</td>
<td>10,000</td>
<td>1,750</td>
</tr>
<tr>
<td>Total</td>
<td>109,000</td>
<td>75,750</td>
</tr>
</tbody>
</table>

The economic efficiency analysis (above) describes the success of contamination management as illustrated by a manufacturing line in the automotive industry with an output of 3,000 systems/day. Manufacturing is done 260 days/year. A contamination review showed that the cleanliness of the function test stand fluid, the intermediate storage conditions and a machining process had to be optimized.

The next step involved forwarding the cleanliness specifications to the suppliers, who received orientation training and are periodically monitored.

The results of optimization:

- Less tool wear in surface machining
- Longer service life of the machining fluid
- Enhanced effectiveness of the downstream washing processes as less dirt had to be removed thanks to optimized storage and machining
- Longer intervals between changing the washing and flushing fluids, consequently “Saturday shifts” could be dispensed with
- Fewer outages at the test stand, i.e. the system is checked up to 3 times when performance deviations occur. These “idle cycles” were reduced by 90%, thus resulting in increased productivity.
- Drop in warranty and non-warranty courtesy work by 50% as the main reason for the outages turned out to be particulate contamination, which resulted in leakage and imprecise control in the system.
- Shortening of the test stand time.

Unfortunately we were not permitted to publish the detailed data behind these savings. Following from an economic efficiency analysis conducted by the customer in-house, savings of $0.60 per system were achieved.

When expressed in terms of the company’s annual output of 780,000 systems, this translates into savings of: $468,000

This economic efficiency analysis also includes the expenses associated with contamination management (seminars, consulting fees, analysis costs).
In the previous pages we discussed the impacts of particulate contamination on the service life and reliability of hydraulic systems, how the cleanliness of fluids on components can be specified, and how contamination monitoring is performed. Deploying contamination management results in the following tasks for all participants in the production process:

Suppliers: Ensuring the defined as-supplied condition of products. Selecting the packaging of products to be supplied so that no additional contamination occurs during transportation and storage.

System vendors and manufacturers: Careful transportation, handling, storage and unpacking of products. Keep products clean after they are unpacked or after seals/plugs have been removed. Assemble/install the components in a suitably clean environment.

The following example shows how these individual parts can be combined in contamination management.

Description of the Point of Departure
System X has been successfully manufactured and marketed for years. During the past few years, System X has been developed further and a new generation, System Y, created. Y features improved performance properties, is more compact than X, and operates at higher system pressures than X. The result is that System Y is somewhat more sensitive to particulate contamination.

This is reflected in increased performance deviations during function testing. This deviation no longer occurs when Y is passed through the test stand a second or third time. An investigation of the matter has shown that this unwanted behavior is the result of coarse particulate contamination.

The goal of contamination management is now to improve the degree of cleanliness so that this undesirable behavior no longer occurs on the test stand and the associated costs of warranty and non-warranty courtesy work are reduced.

Step 1: Analysis of the Test Fluid
The cleanliness of the test fluid is determined. The analyses show that the test fluid cleanliness upstream of the test item amounts to a cleanliness rating of 22 / 20 / 18 according to ISO 4406, the largest metallic particles are 400 µm in size, and the largest fibers measure 3,000 µm.

Step 2: Optimizing the Function Test Stand
By additionally integrating bypass microfiltration, which maintains test fluid cleanliness at 15 / 13 / 10, 95% of the performance deviations can be prevented. This also results in a drop in warranty and non-warranty courtesy work.

Step 3: Lowering the Filter Costs at the Test Stands
By performing a contamination monitoring audit, it might be determined a large amount of particulate contamination is being transported into the system by the manufacturing processes and sourced components. This particulate contamination has to be removed from the system at the function test stand, which functions here as the last washing operation. This results in costs that could otherwise be avoided.

A concept is developed in which the washing and machining processes and intermediate storage are optimized.

A cleanliness specification along with a test plan for system fluids is drafted. This specification is forwarded to external as well as internal suppliers and the components supplied with a defined, constant cleanliness.

Step 4: Integrating Particle Counting in Quality Assurance
A particle sensor is integrated in the function test stand for the purpose of continuous quality control of the as-supplied quality of System Y. A limit is defined for the maximum contamination of the test fluid in the return line. Intervention can be done immediately if this value is exceeded, thus ensuring that no contaminated systems leave the factory. Random sampling is done to check the supplier quality and non-conformant components returned to suppliers or washed in-house at the supplier’s expense.

Step 5: Economic Efficiency Analysis
Contamination management started off with analyzing the costs associated with warranty and non-warranty courtesy work as the result of increased malfunction at the test stands. These costs are reanalyzed after optimization and compared. The savings achieved through optimization are briefly described in Economic Efficiency Analysis. The cost savings in that case amounted to ca. €468,000/year. This optimization process lasted ca. 2 years.

Step 6: Documentation and New Projects
The contamination management findings are collected in a database and used in the development of new systems. The defined maximum residual dirt content becomes standard in new systems in the same way that dimensions, surface grades and tolerances have been. This residual dirt content is primarily in reference to the specification that applies to System Y.

The specification is adapted in keeping with the experience gained with the prototypes. Cleanliness and cleaning costs are primarily determined by the design of new systems.