



Translation of the original user’s manual

Introduction

Dear customer,
thank you for the confidence you have shown in the Extol® brand by purchasing this product. This product has been tested for reliability, safety and quality according to the prescribed norms and regulations of the European Union.

Contact our customer and consulting centre for any questions at:

extol.eu

Manufacturer: Madal Bal a. s., Průmyslová zóna Příluky 244, 76001 Zlín, Czech Republic.
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EN

TECHNICAL SPECIFICATIONS OF RESPIRATORS

Model/part number of the respirator	8856716	8856724	8856738
Filter designation	FFP1 NR	FFP2 NR D	FFP3 NR D
Layer of active carbon			•
Exhalation valve	•	•	•
Type of respirator	Folding	Formed	Formed

Table 1

Explanations for table 1:

The symbol “•” means YES; absence of the symbol in the table means NO.

The filter class (efficiency) FFP1, FFP2, FFP3: filter filtration efficiency classification according to the harmonised norm EN 149 with further specifications provided in table 2.

NR: Respirator intended for single disposable use (it cannot be cleaned for repeat use).

D: The respirator passed the dolomite dust clogging test according to EN 149+A1.

Layer of active carbon: Respirators with a layer of active carbon are suitable for cases where unpleasant non-toxic odours are present, or for the occurrence of ozone from road traffic, smog and furthermore in cases recommended according to table 2.

Exhalation valve: This is a one-way flap that closes upon inhalation, i.e. inhaled air is filtered through the material of the respirator, and opens upon exhalation. The exhalation valve extracts exhaled air out of the respirator, which increases comfort when the respirator is used, since it reduces the heating of the wearer’s face, particularly in higher ambient temperatures or during physically demanding work, which increases the breathing frequency.

Formed/folding type:

The formed type of respirator has a permanent form (body) and increases the comfort of the user during more demanding work, where the respirator retains its shape during intensive inhalation or when worn in windy weather.

The folding type of respirator has a soft form and thanks to this can be folded for easier carrying, e.g. in the pocket of clothing or in a handbag.

Material of the respirator: polypropylene (PP)

USING THE RESPIRATOR ACCORDING TO THE FILTRATION EFFICIENCY CLASS FFP1, FFP2 AND FFP3

- Respirators are intended for protection against the inhalation of solid particulates and aerosols based on the filtration efficiency class FFP1, FFP2 and FFP3, the national assigned protection factor APF¹⁾ and permitted exposure limits for given harmful substances (see below). Classification of filtration classes FFP1, FFP2 and FFP3 according to efficiency is given by the harmonised norm EN 149.

Filter class (efficiency) – use according to NPK-P/PEL	Example of material particulate types
FFP1 Filtration efficiency of respirator material min. 80% Total penetration through respirator: max. 22% Against non-toxic dust and aerosols up to a concentration of 4 times the permitted exposure limit (PEL) for the given harmful substance.	Cement ²⁾ , limestone, cinder, cereal grain dust, dust from abrasives, plaster ²⁾
FFP2 Filtration efficiency of respirator material min. 94% Total penetration through respirator: max. 8% Against dust with predominantly irritating effects and aerosols up to a concentration of 10 times the permitted exposure limit (PEL) for the given harmful substance.	- Cotton, flax, hemp, silk, synthetic textile fibres, (textile industry) - Feathers, animal fur, pollen, mites (allergens) - Flour, tobacco, tea, coffee, spices, dust from dried herbs (food processing) - Fibreglass, glass, resin, concrete, plaster, soft wood, old paint coats, rust (grinding building materials) - Smog and ground-level ozone from road traffic (environment)+layer of active carbon - Unpleasant non-toxic odours + layer of active carbon - Attic dust, soot (cleaning – soiled with standard dust) - Coal, stone (mining activity)
FFP3 Filtration efficiency of respirator material min. 99% Total penetration through respirator: max. 2% Against toxic particulates, viruses, spores, bacteria, radio active dust up to a concentration of 30 times the permitted exposure limit (PEL) for the given harmful substance.	- Viruses³⁾, bacteria, mould spores (harmful biological agents) - Machining hard wood (e.g. oak, beech) - Mineral and glass insulating wool - Welding of stainless steel, aluminium - Welding of steel, zinc - Welding fumes + layer of active carbon - Handling of asbestos and lead

Table 2

Explanations for table 2

ATTENTION

- Layer of active carbon in the respirator (if included): Captures unpleasant non-toxic odours and gases.
- The total penetration through the respirator includes penetration along the contact line of the respirator with the face, penetration through the material of the filter and penetration through the exhalation valve, if the respirator is equipped with one.

¹⁾ Norm EN 529:2005 for recommendations regarding the selection of protective equipment for respiratory organs with effective protection according to class FFP1, FFP2 and FFP3 are based on a so-called nominal protection factor NPf, which is for class FFP1: 4, for class FFP2: 12 and for class FFP3: 50 and which is specified by many different respirator brands, however this factor NPf is specified only from the max. permitted penetration through the respirator given by norm EN 149+A1, according to the formula: 100/x% (total penetration as a percentage), which however does not take into account the toxicity of the substance and its permitted exposure limit PEL according to nation legislation (in CZ according to NV. 361/2007 Coll as amended) and which for this reason varies in several countries precisely as a result of the requirements of national legislation, which is not harmonised. Table 3 specifies the mentioned (national) assigned protection factors (APF) for the individual respirator classes specified in the mentioned countries. For the assessment of the minimum level of required protection, it is necessary to calculate the minimum required protection factor as a ratio: of the concentration of the specific harmful substance outside the contact part / permitted concentration of harmful substances inside the contact part (permitted exposure limit) and to compare this with the national assigned protection factor APF in the given country, in which the respirator is used, and based on this, select the respirator class FFP1, FFP2 or FFP3 (see EN 529).

Filter class EN 149	Assigned protection factor APF used in the listed countries ¹⁾				
	FIN	D	I	S	UK
FFP1	4	4	4	4	4
FFP2	10	10	10	10	10
FFP3	20	30	30	20	20

²⁾ for CZ, the APF factors are specified in table 2, i.e. for FFP1: 4; FFP2: 10; FFP3: 30 Table 3

²⁾ At higher concentration of substance, select FFP2

ATTENTION

- It is essential to understand that not even respirators with the highest filtration efficiency of class FFP3 can entirely eliminate the risk of infection by damaging biological agents such as viruses and bacteria, because these infectious particles that are present in the air come into contact with eyes, ears, fine skin injuries and may enter into the organism through these pathways, and likewise they adhere to clothing with which an unprotected person comes into contact. For effective protection against damaging biological agents, the user must, for the entire duration of the exposure, wear a special protective suit with a high level of protection against biological hazards including protection of the entire head with a separate supply of uncontaminated oxygen and prior to undressing must undergo a thorough decontamination process! Other interesting interrelationships with regards to the filtration efficiency of FFP3 respirators in relation to damaging biological agents. The filtration effectiveness of FFP3 class materials is not 100% (“only” 99%) and the total penetration through the respirator is not zero, but may be up to 2% (according to norm EN 149+A1 at least 8 out of 10 tested samples must meet the specified filtration effectiveness values and the total penetration through the filtration face mask, i.e. not all 10 samples).
- Furthermore, it is necessary to stress that the filtration efficiency of the material is tested using a test aerosol with a flow rate of 95 l/min through the respirator with an aerosol particulate diameter in the range from 20 nm to 2000 nm, whilst the mean size of the aerosol particulates is 600 nm and the diameter of viral particulates, depending on the type of virus, is in the range from 23 nm (rhinovirus cold virus) - 63 nm (rubella/German measles togavirus) - 100 nm (flu virus 1 - orthomyxovirus) - 112 nm (coronavirus (SARS2)), and it thus follows that the respirator with class FFP3 cannot provide absolute protection against viruses, nevertheless can significantly reduce the risk of infection. Note: the size of bacteria is significantly larger and is in the range from approx. 1000 nm length × approx 600 nm - 1500 nm width for rod-shaped bacteria.

ATTENTION

- The here provided examples of harmful substances in table 2 are general recommendations and for the protection of the user it is necessary to know the exact concentration of all harmful substance in the air and the permitted exposure limits for the specific harmful substance according to the legislation of the country in which the respirator is used. Assessment of the minimum required level of protection must relate to both the concentration of harmful substances with the highest toxicity as well as the concentrations of other harmful substance in the environment with lower toxicity.

PEL: The permitted exposure limit to a chemical substance, dust, aerosol in the air, to which an employee may be exposed during an eight-hour or shorter weekly work shift, see. NV. 361/2007 Coll. as subsequently amended.

- The requirement for the minimum filtration efficiency of the respirator material and the total penetration through the respirator for the given class FFP1; FFP2 and FFP3 is specified by harmonisation norm EN 149+A1.
- Total penetration includes parts: penetration through the sealing line part of the half face mask, penetration through the material of the filter and penetration through the exhalation valve (if part of the respirator).

SAFETY WARNINGS

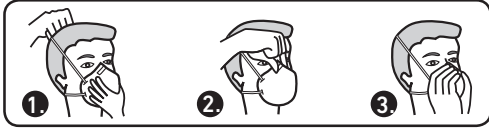
FOR THE USE OF RESPIRATORS

- The respirator does not protect against gases and lower oxygen content in air.
- The respirator must not be used for protection in an environment where the composition and concentration of all harmful substances in the air are unknown.
- The respirator must not be used for longer than the duration of one work shift, thereafter it must be disposed off! According to EN 149+A1 a respirator used for a duration of no more than a single work shift is considered to be disposable. Its lifetime is, however, dependent on the concentration of particulates in the workplace and the activity of the user, i.e. at a higher particulate concentration and higher breathing frequency of the user it must be replaced with a new one sooner than the duration of a single work shift.
- The respirator must be replaced when dust penetrates through the respirator and internal soiling, odour or other evidence of penetration of harmful substances through the respirator is identified on the inside of the respirator; furthermore also when it is damaged or breathing resistance increases significantly (breathing becomes difficult) or when the user has a feeling of insufficient oxygen supply.
- The respirator is not intended to be cleaned and disinfected for repeat use.
- The respirator is not intended to be used in an environment where there is a risk of explosion due to the risk of the ignition of flammable fumes resulting from static electricity from the dust on the respirator.
- Tight seal requirements will most probably not be met if the user has whiskers on the contact surface of the sealing line of the respirator.
- The respirator should not be used by persons whose whiskers, facial shape or scars prevent proper tight contact of the respirator with the face.
- Not following the instructions and warnings may reduce the efficiency of the respirator and may result in damage to health.

PROCEDURE FOR FITTING THE RESPIRATOR ON THE FACE:

ATTENTION

- To ensure that the respirator provides protection; it is necessary to fit it correctly on the face because an incorrectly fitted or located respirator will significantly reduce the level of protection!
- Tight seal requirements will not be met if the user has whiskers or hair on the contact surface of the sealing line.
- Prior to fitting the respirator, check that it is complete and that it has not been damaged mechanically. Do not use respirators that have been used for longer than one work shift, respirators with increased breathing resistance and clogging, or that are past the expiry date shown on the packaging of the respirator.



- Position the top and bottom fastening rubber bands into the positions according to fig. 1, so that the top and bottom parts of the respirator are seated as tightly as possible in contact with the face.
- With your fingers, thoroughly squeeze the nose strip against the nose and form it along the contour of the nose so that the respirator holds as best as possible on the face and seals along the contact line (fig. 2).
- Check the respirator for leaks along the contact lines with the face by forcefully and intensively breathing in. If the respirator is not sealing and air is passing around the nose or elsewhere along the sealing line, it is necessary to readjust the respirator by moving it to a more suitable position, changing the shape of the nose strip or moving the fastening rubber bands to a more suitable position, see fig. 3.

ATTENTION

- In the event of increased breathing resistance, replace the respirator for a new one.

STORAGE

- The respirator must be stored in an environment with a temperature of +5° to +38 °C at a relative humidity of < 70%.
- The lifetime (expiry date) is provided on the sales packaging (see respective pictogram in table 4) under the condition that the specified storage conditions are adhered to.

MEANINGS OF MARKINGS ON THE RESPIRATOR AND SMALLEST SALES PACKAGING

EXTOL® P R E M I U M	Manufacturer's trade mark. The address of the manufacturer is provided on the product in the form: Made by Madal Bal a.s., Prům. zóna Příluky 244, CZ-76001 Zlín		
EN 149:2001 + A1:2009	Harmonisation norm defining the requirements for filtration half face masks for protection against particulates; norm's year of issue.		
CE	It meets all the respective provisions of the harmonisation legal regulation for personal protective equipment (EU) 2016/425.		
CE 2797	Number of the notified entity that performed the EU assessment of the filtration half face mask type tested according to EN 149+A1:2009 and that performs supervision of production according module D.		
FFPX NR D	"FFP"- "Filtering Facepiece Particles" - "Face filtration mask against solid particulates". "X" – filter class 1; 2 or 3 according to filtration efficiency. "NR" – designed for use for the duration of a single shift only, then the respirator must be disposed of. "D" – The respirator passed the dolomite dust clogging test according to EN 149+A1 (applies for only the models specified in table 1).		
Expiry date M/YYYY	Final lifetime date of the respirator (expiry date).		Read the user's manual before using the respirator.
+5 °C – +38 °C	Storage temperature range for the respirator.	<70 %	Maximum humidity for the storage of the respirator.

Table 4

EU DECLARATION OF CONFORMITY

Subjects of declaration - Product identification:

Respirators: Extol® Premium 8856716 (FFP1 NR);

Extol® Premium 8856724 (FFP2 NR D); Extol® Premium 8856738 (FFP3 NR D).

Manufacturer Madal Bal a.s.- Bartošova 40/3, CZ-760 01 Zlín - Corp. ID NO:49433717

hereby declares, that the described products listed above are in conformity with the harmonisation legal regulation of the European Union: (EU) 2016/425. This declaration is issued under the exclusive responsibility of the manufacturer.

Harmonisation norm (including its amendments, if any exist), which was used in the assessment of conformity and on the basis of which the declaration of conformity is issued: EN 149:2001+A1:2009

Notified Body No. 2797; BSI Group The Netherlands B.V., John M. Keynesplein 9, 1066 EP Amsterdam, Holland, carried out an EU type - examination according to module B and issued certificate no.: CE 837751 dated 02.01.2026. The aforementioned personal protection products (respirators) are also subject to the procedure for the assessment of conformity with the type based on ensuring the quality of the production process according to module D (certificate no.: CE 737355) under the supervision of the aforementioned notified body no.: 2797 BSI Group The Netherlands B.V.
Place and date of issue of EU Declaration of Conformity: Zlín, 02.01.2026
On behalf of Madal Bal, a.s.:

Martin Senkýř, Member of the Company's Board of Directors