INTRA

| | I.1. Consignor | | | | I.2. IMSOC reference | | I.2.a. Local refere | ence | |
|---|--|-------------------------|------------------------------|----------------------------|--|---------------|---------------------|-----------------|--|
| | Name Address | | | | | | I.3. Central Comp | etent Authority | |
| | Country | | ISO Code | | | | I.4. Local Compet | tent Authority | |
| ent | I.5. Consignee | | | | I.6. Operator conducting assembly operations independently of an establishment | | | dently of an | |
| nu | Name Address | | | | Name | | | | |
| sig | Country | | ISO Code | | Address | | | | |
| Ö | | | | | Approval Number | | | | |
| $\overline{\mathbf{f}}$ | | | | | Country | | ISO Code | | |
| Part I: Description of consignment | I.7. Country of ori | gin | ISO Cod | e | I.9. Country of destination | on | | ISO Code | |
| crip | I.8. Region of origin Code I.11. Place of dispatch | | | | I.10. Region of destination Code | | | | |
| Des | | | | I.12. Place of destination | | | | | |
| Γ: | Name | | | | Name | | | | |
| art | Address Approval | | | | Approval | Approval | | | |
| ľ | Number | | 100.0.1. | | Number | | 100 0 1 | | |
| | Country | | ISO Code | | Country | | ISO Code | | |
| | I.13. Place of loadi | ing | | | I.14. Date and time of de | parture | | | |
| | Name Address | | | | | | | | |
| | Approval Number | | | | | | | | |
| | Number Country | | ISO Code | | | | | | |
| | | | 100 code | | | | | | |
| | I.15. Means of Tra Mode | nsport International | Identification | | I.16. Transporter Name | | | | |
| | Wode | transport document | lacitification | | Address | | | | |
| | | document | | | Approval | | | | |
| | | | | | Nûmber Country | | ISO Code | | |
| | | | | | I.17. Accompanying documents | | | | |
| | | | | | Document Type | | | | |
| | | | | | Accompanying document | | | | |
| | | | | | reference Date of Issue Country Place of issue | | | | |
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| | | | | | | | | | |
| | I.18. Transport con Chilled □ | nditions | Ambient □ | | | Frozen 🗆 | | | |
| | | /0.122 | | | | | | | |
| | I.19. Container No | o / Seal No | | | | | | | |
| | I.20. Certified as | | | | | | | | |
| | Germinal product | S L | | | | | | | |
| | I.21. For transit th | rough a third coun | try | | | | | | |
| | Third country | | | | ISO Code | | | | |
| | Exit point | | | | BCP code | | | | |
| ŀ | Entry point I.22. For transit through Member State(s) | | | BCP code I.23. For export | | | | | |
| | Member State | noagn meniber old | ISO Code | | Third country | | ISO Code | | |
| | MEHINEL STATE | | 130 Code | | Exit point | | BCP code | | |
| ı | I.24. Estimated jou | ırney time | | | I.25. Journey Log | | | | |
| | I.26. Total number | r of packages | I.27. Total qua | intity | | I.28. Total g | ross weight | | |
| | I.30. Description o | of consignment | 1 | | | 1 | | | |
| | 1. 05 PRODUCTS O | OF ANIMAL ORIGIN | , NOT ELSEWHERE SPECIFI | ED OR IN | CLUDED | | | | |
| | 0511 Animal pr 051110 Bovin | | re specified or included; de | ad anima | ls of Chapter 1 or 3, unfit | for human o | consumption | | |
| | 05111000 E | Bovine semen | | | | | | | |

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| _ | NION | | | | | INTRA |
|------------------------------------|---------------|-----------------------|---------------------------------|--------------------------------|---------------------|-------|
| | #1. Commodity | Identification Number | Quantity | Nature of commodity | Identification Mark | |
| | Species | Package count | Date of collection / production | Plant / Establishment / Centre | | |
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| Part I: Description of consignment | | | | | | |
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| -01 | NION | | | | | 2024/1044 (202 | 11/403) 1 | Model BOV-SEM-A-IN I | .KA | | |
|--|-----------------|------------------------|---------------|---|---|---|---------------------------------|--|-------------|--|--|
| | II. Health info | ormation | | | | | | | | | |
| | I, the unde | ersigned off | icial veterii | narian, hereb | ا y certify that | <u> </u> | | | | | |
| | II.1. | • | | animals desci | | | | | | | |
| | II.1.1. | | _ | ocessed and s a register by | | _ | semen co | ollection centre (1) which | is | | |
| lo | II.1.2. | | _ | | | - | | ollection centre (1) which | | | |
| ficati | | | | | sibilities, operationa hission Delegated Re | | | | | | |
| II.1.2. has been collected, processed and stored, and dispatched from the semen collection centre of complies with requirements as regards responsibilities, operational procedures, facilities at equipment set out in Part 1 of Annex I to Commission Delegated Regulation (EU) 2020/686; (2) either (2) either (3) is dispatched from a semen collection centre or a zone not subject to movement affecting bovine animals and established for reasons of listed diseases relevant affecting bovine animals and established for reasons of listed diseases relevant for those species, or restrictions do not apply to this semen because it was collected before the restrictions do not apply to this semen because it was collected before the restrictions do not apply to this semen because it was collected before the restrictions do not apply to this semen because it was collected before the restrictions do not apply to this semen because it was collected before the restrictions do not apply to this semen because it was collected before the restrictions do not apply to this semen because it was collected before the restrictions do not apply to this semen because it was collected before the restrictions do not apply to this semen because it was collected before the restrictions do not apply to this semen because it was collected before the restrictions do not apply to this semen because it was collected before the restrictions do not apply to this semen because it was collected before the restrictions do not apply to this semen because it was collected before the restrictions do not apply to this semen because it was collected before the restrictions do not apply to this semen because it was collected before the restrictions do not apply to this semen because it was collected before the restrictions do not apply to this semen because it was collected before the restrictions do not apply to this semen because it was collected before the restrictions do not apply to this semen because it was collected before the restrictions do not ap | | | | | | | | | re | | |
| | (2) or | ○ [II.1.3. | affecting k | ned from a ser povine animal t restrictions l | ls and establ | ished for | | movement restrictions at derogations from | | | |
| | (2) | ☐ [it comp | olies with th | ie requiremer | nts set out in | (4);]] | | | | | |
| | (2) | | particular, | | (5).]] | | | | | | |
| | II.2. | The semer animals w | | in Part I is in | tended for a | rtificial reproduction | n and wa | s obtained from donor | | | |
| | | II.2.1. | | | | e birth in the Union, or have entered the Union in for entry into the Union; | | | | | |
| | | II.2.2. | establishn | nents in a Mei | mber State o | | om estab | red to in point II.2.6, from lishments under official y, or a zone thereof: | 1 | | |
| | | | II.2.2.1. | 10-km radiu | s centred on | re foot and mouth disease has not been reported within a name the establishment for at least 30 days and in which foot not been reported during at least 3 months, and: | | | | | |
| | | (2) | o either | [the donor a | nimals were | not vaccinated agai | nst foot a | and mouth disease;] | | | |
| | | (2) | o or | months prio immediately minimum of | or to the date or prior to the f five straws) is submitted | of collection of the s date of collection of of each quantity of | semen bu the sem semen ta | mouth disease during the at not during the last 30 den, and 5 % (with a aken from the donor anime and mouth disease with | ays nals | | |
| | | | II.2.2.2. | caprae and l | M. tuberculo | | nimals h | s complex (M. bovis, M. nave never been kept tus; | | | |
| | | | | | | | | sis and B. suis, and the do blishment of a lower heal | | | |
| | | (2) | o either | [II.2.2.4. | | enzootic bovine leukosis, and the donor animals have en kept previously in any establishment of a lower health | | | | | |
| | | (2) | o or | [II.2.2.4. | younger th have been | rom enzootic bovine leukosis and the donor animals are than 2 years of age and have been produced by dams whin subjected, with negative results, to a serological test for bovine leukosis after removal of the animal from the dam | | | | | |
| | | (2) | o or | [II.2.2.4. | reached the | | have bee | and the donor animals ha en subjected, with a negat povine leukosis;] | - 1 | | |
| | | (2) | o either | [II.2.2.5. | free from i | nfectious bovine rhi | notrache | eitis/infectious pustular | | | |

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| | NION | | | | 2024/1044 (2021/403) | Model BOV-SLM-A-INTRA | | | |
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| | II. Health information | | | | | | | | |
| | | | | | itis, and the donor animals lin any establishment of a lo | | | | |
| cation | (2) | o or | [II.2.2.5. | vulvovagin | m infectious bovine rhinotr itis and the donor animals h sult, to a serological test (wh | ave been subjected, with a | | | |
| tificat | (2) | II.2.2.6. | in which su days, and: | rra (Trypano | soma evansi) has not been r | eported during the last 30 | | | |
| Cer | | o either | [surra has not been reported in the establishments during the last 2 years.] | | | | | | |
| Part II: Certification | (2) | ∘ or | following the remained used animals have in the estab diagnostic regulation of the state of the | e been subjected to a test fo ided for in Part 3 of Annex I , carried out, with negative the date on which the infec | stablishments have e on which the infected , and the remaining animals r surra with one of the to Commission Delegated results, on samples taken at | | | | |
| | II.2.3. | | | | igns of transmissible anima tre and on the date of collec | diseases on the date of their tion of the semen; | | | |
| | II.2.4. | are indivi (EU) 2019 | | fied as provid | ed for in Article 38 of Comn | nission Delegated Regulation | | | |
| | II.2.5. | for at leas period: | st 30 days pric | or to the date | of first collection of the sem | en and during the collection | | | |
| | | II.2.5.1. | restrictions with rinder | established d pest virus, in imonia or lum | ents situated in a zone not sulue to the occurrence of foot fection with Rift Valley feve apy skin disease, or of an en | and mouth disease, infection virus, contagious bovine | | | |
| | | II.2.5.2. | melitensis a bovis, M. ca evansi), enz pustular vu haemorrha | and B. suis, in prae and M. to cootic bovine lvovaginitis, l gic disease vi | ablishment where infection fection with Mycobacterium uberculosis), rabies, anthra leukosis, infectious bovine rovine viral diarrhoea, infectus, infection with bluetong acteriosis and trichomonosi | tuberculosis complex (M. x, surra (Trypanosoma hinotracheitis/infectious ction with epizootic ue virus (serotypes 1-24), | | | |
| | | II.2.5.3. | to movemen | nt restrictions om establish | animals from establishmen s due to the occurrence of di ments which do not meet th | | | | |
| | | II.2.5.4. | were not us | sed for natura | l breeding; | | | | |
| | II.2.6. have been subjected to a quara where only other cloven-hoofe which on the date of admission with the following conditions: | | en-hoofed ani dmission of th ditions: | mals with at least the same in the same in the donor animals to the sem | health status were present, en collection centre complied | | | | |
| | | | | ferred to in po | oint II.2.5.1 or it was under o | strictions established due to lerogation as referred to in | | | |
| | | II.2.6.2. | none of the days; | diseases refe | rred to in point II.2.5.2 has l | peen reported for at least 30 | | | |
| | | II.2.6.3. | | | where foot and mouth disentred on the quarantine acc | - 1 | | | |

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| | | | 2021/1011 (2021/100) Model Bo V OLM 11 INVITED |
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| II. Health in | formation | | |
| | | II.2.6.4. | it has had no outbreak of foot and mouth disease reported during at least 3 months prior to the date of admission of the donor animals into the semen collection centre; |
| | II.2.7. | were kept | in the semen collection centre: |
| Part II: Certification (2)(2) (2)(7) (2)(7) | | II.2.7.1. | which was situated in a zone not subject to movement restrictions established due to diseases referred to in point II.2.5.1 or it was under derogation as referred to in point II.1.3, if applicable; |
| Certif | | II.2.7.2. | where none of the diseases referred to in point II.2.5.2 has been reported for at least 30 days prior to the date of collection of the semen, and: |
| (2)(6) | | | ☐ [at least 30 days following the date of the collection;] |
| (2)(7) | | | \square [until the date of dispatch of the consignment of semen to another Member State;] |
| | | II.2.7.3. | situated in an area where foot and mouth disease has not been reported within a 10-km radius centred on the semen collection centre for at least 30 days; |
| | II.2.8. | | ith at least one of the following conditions as regards infection with bluetongue otypes 1-24): |
| (2) | □ either | [II.2.8.1. | they have been kept for at least 60 days prior to and during collection of the semen in a Member State or zone thereof free from infection with bluetongue virus (serotypes 1-24) where no case of infection with bluetongue virus (serotypes 1-24) has been confirmed during the preceding 24 months in the targeted animal population;] |
| (2) | □ or | [II.2.8.2. | they have been kept in a seasonally disease-free zone, during the seasonally disease-free period, for at least 60 days prior to and during collection of the semen, in a Member State or zone thereof with an approved eradication programme against infection with bluetongue virus (serotypes 1-24);] |
| (2) | □or | [II.2.8.3. | they have been kept in a seasonally disease-free zone, during the seasonally disease-free period, for at least 60 days prior to and during collection of the semen, in a Member State or zone thereof where the competent authority of the place of origin of the consignment of semen has obtained the prior written consent of the competent authority of the Member State of destination to the conditions for establishment of that seasonally disease-free zone and to accept the consignment of semen;] |
| (2) | □ and/or | [II.2.8.4. | they have been kept in a vector protected establishment for at least 60 days prior to and during collection of the semen;] |
| (2) | □ and/or | [II.2.8.5. | they have been subjected to a serological test to detect antibodies to the bluetongue virus serotypes 1-24, with negative results, between 28 and 60 days from the date of each collection of the semen;] |
| (2) | □ and/or | [II.2.8.6. | they have been subjected to an agent identification test for bluetongue virus (serotypes 1-24), with negative results, on blood samples taken at commencement and final collection of the semen and during collection of the semen at intervals of at least every 7 days, in the case of the virus isolation test, or of at least every 28 days, in the case of PCR;] |
| | II.2.9. | | ith at least one of the following conditions as regards infection with epizootic nagic disease virus (EHDV): |
| (2) | □ either | [II.2.9.1. | they have been kept for at least 60 days prior to and during collection of the semen in a Member State or zone thereof where EHDV has not been reported for at least 2 years within a radius of 150 km of the establishment;] |
| (2) | □ or | [II.2.9.2. | they have been kept in a seasonally disease-free zone, during the seasonally disease-free period, for at least 60 days prior to and during collection of the semen;] |
| (2) | □ and/or | [II.2.9.3. | they have been kept in a vector protected establishment for at least 60 days prior to and during collection of the semen;] |
| (2) and/o | r 🗆 | were resid | dent in the Member State in which according to official findings the following |

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| | NION | | | | | 2024/1044 (2021/403) | WIGHEI BOV-SLWI-A-INTRA | | | | | |
|------------------------|-------------------------------------|------------|--|---|--|---|--|--|--|--|--|--|
| | II. Health info | ormation | | | | | | | | | | |
| | | [II.2.9.4. | serotypes of EHDV exist: and have been subjected with negative results in case to the following tests carried out in an official laboratory: | | | | | | | | | |
| u | (2) | | □ either | [II.2.9.4.1. a serological test to detect antibodies to EHDV, with nega at least every 60 days throughout the collection period at 28 and 60 days from the date of the final collection of the | | | | | | | | |
| Part II: Certification | (2) | | □ and/or | [II.2.9.4.2. | samples ta semen and every 7 day | ken at the commencement ar l during the collection of the s | | | | | | |
| Part I | | II.2.10. | prior to th negative r II.2.10.5.2, | have been subjected to the following tests, carried out on blood samples taken within 30 days prior to the date of commencement of the quarantine referred to in point II.2.6, with negative results, except for the bovine viral diarrhoea antibody test referred to in point II.2.10.5.2, required in accordance with Part 1, Chapter I, point 1(b), of Annex II to Delegated Regulation (EU) 2020/686: | | | | | | | | |
| | | | II.2.10.1. | M. tubercul | osis), an intr | bacterium tuberculosis comp adermal tuberculin test refer gulation (EU) 2020/688; | _ | | | | | |
| | | | II.2.10.2. | | | lla abortus, B. melitensis and int 1, of Annex I to Delegated | • | | | | | |
| | | (2)(8) | □ [II.2.10.3. | | | vine leukosis, a serological test referred to in Part 4, gated Regulation (EU) 2020/688;] | | | | | | |
| | | | II.2.10.4. | serological t | test (whole v ment free fr | inotracheitis/infectious pustu irus) on a blood sample if the om infectious bovine rhinotr | animals do not come from | | | | | |
| | | | II.2.10.5. | for bovine v | riral diarrho | ea: | | | | | | |
| | | | | II.2.10.5.1. | a virus isol antigen, ar | lation test, a test for virus ger nd | ome or a test for virus | | | | | |
| | | | | II.2.10.5.2. | a serologic antibodies | al test to determine the prese; | ence or absence of | | | | | |
| | | II.2.11. | days in the commence the bovine | een subjected to the following tests, carried out on samples taken at least 21 the case of the tests referred to in points II.2.11.4 and II.2.11.5, after the encement of the quarantine referred to in point II.2.6, with negative results, rine viral diarrhoea antibody test referred to in point II.2.11.3.2, required in ance with Part 1, Chapter I, point 1(c), of Annex II to Delegated Regulation (c). | | | | | | | | |
| | | | II.2.11.1. | | | lla abortus, B. melitensis and int 1, of Annex I to Delegated | • | | | | | |
| | | | II.2.11.2. | | | inotracheitis/infectious pustu irus) on a blood sample; | lar vulvovaginitis, a | | | | | |
| | II.2.11.3. for bovine viral diarrho | | ea: | | | | | | | | | |
| | | | | II.2.11.3.1. a virus isc antigen, a | | olation test, a test for virus genome or a test for virus nd | | | | | | |
| | | | | II.2.11.3.2. a serologic antibodies | | al test to determine the prese; | ence or absence of | | | | | |
| | | | II.2.11.4. | for bovine genital campy | | ylobacteriosis (Campylobacte | r fetus ssp. venerealis): | | | | | |
| | | (2) | □ either | [II.2.11.4.1. | preputial s kept since | st carried out on a sample of a specimen, in the case of anim that age in a single sex group ior to the quarantine referre | als less than 6 months old or without contact with | | | | | |
| | | (2) | \square and/or | [II.2.11.4.2. | tests carrie | ed out on samples of artificial | vagina washings or | | | | | |

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| | II. Health info | ormation | | | | | | | | |
|--|--|--------------|--|---|---------------|-------|---|-----------|----------------------------|---|
| | | | | | preputial s | spec | imens | taken or | three occ | asions at intervals of at least |
| | | | | | 7 days;] | орсс | ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,, | tuncii oi | r an co occ | uorono at mitor valo or at roust |
| | | | II.2.11.5. | for trichomo | onosis (Trich | hom | ionas f | oetus): | | |
| ication | | (2) | [II.2.11.5.1. | [II.2.11.5.1. a single test carried out on a sample of preputial specimen, in the case of animals less than 6 months old or kept since that age in a single sex group without contact with females prior to the quarantine referred to in point II.2.6.;] | | | | | | |
| ertifi | (2) and/or [II.2.11.5.2. tests car interval: | | | | | | | | specimen | s taken on three occasions at |
| Part II: Certification | | ts, required | emen collection centre, at least once a year, to the following s, required in accordance with Part 1, Chapter I, point 2, of Annex II (EU) 2020/686: | | | | | | | |
| | | | II.2.12.1. | 1. for infection with Mycobacterium tuberculosis complex (M. bovis, M. caprae and M. tuberculosis), an intradermal tuberculin test referred to in Part 2, point 1, of Annex I to Delegated Regulation (EU) 2020/688; | | | | | | |
| | | | II.2.12.2. | | | | | | | B. suis, a serological test Regulation (EU) 2020/688; |
| | | | II.2.12.3. | for enzootic Annex I to D | | | | | | ed to in Part 4, point (a), of |
| | | | II.2.12.4. | for infectious serological t | | | | | - | llar vulvovaginitis, a |
| | | (2)(9) | □ [II.2.12.5. | for bovine viral diarrhoea, a serological test for detection of an antibody;] 3.12.5. | | | | | | |
| | | (2)(10) | [II.2.12.6. | ☐ for boving test on a san | | | | | Campyloba | cter fetus ssp. venerealis), a |
| | | (2)(10) | □ [II.2.12.7. | for trichomo specimen;] | onosis (Trich | hom | ionas f | oetus), a | test on a sa | ample of preputial |
| | II.3. | The semen | n described | in Part I: | | | | | | |
| | | II.3.1. | | | | | | | | mal health requirements set n (EU) 2020/686; |
| II.3.2. is placed in straws or other packages on which the mark is applied requirements provided for in Article 10 of Delegated Regulation (EU is indicated in box I.30; | | | | | | | | | | |
| | | II.3.3. | is transported in a container which: | | | | | | | |
| | | | II.3.3.1. | centre unde | r responsibi | ility | of the | centre v | eterinariaı | ch from the semen collection n, or by an official ated in box I.19; |
| | | | II.3.3.2. | has been cle container; | eaned and ei | ithe | r disin | fected or | sterilised | before use, or is single-use |
| | | (2)(6) | □ [II.3.3.3. | has been fill other produ | | a cry | ogenio | c agent w | hich has n | ot been previously used for |
| | (2) | □ [II.4. | Where an antibiotic or a mixture of antibiotics was added to the semen described in Part I: | | | | | | | |
| II.4.1. The following antibiotic or mixture of antibiotics has been added to the semen after fin dilution, or is contained in the used semen diluents:(11). | | | | | | | | been adde | d to the semen after final | |
| | | | | | | | | | | |
| | II.4.2. | was kept a | at a tempera | | st 5°C for no | ot le | ss thai | n 45 mini | | freezing, the diluted semen der a time-temperature |
| | Notes | | | | | | | | | |

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II. Health information

In accordance with the Agreement on the withdrawal of the United Kingdom of Great Britain and Northern Ireland from the European Union and the European Atomic Energy Community, and in particular Article 5(4) of the Protocol on Ireland/Northern Ireland in conjunction with Annex 2 to that Protocol, references to the Union in this animal health certificate include the United Kingdom in respect of Northern Ireland.

This animal health certificate shall be completed in accordance with the notes for the completion of certificates provided for in Chapter 2 of Annex I to Commission Implementing Regulation (EU) 2020/2235.

Part I:

Certification Box reference

"Place of dispatch": Indicate the unique approval number and the name and address of the semen collection centre of dispatch of the consignment of semen.

I.11:

Box reference

"Place of destination": Indicate the address and unique registration or approval number of the establishment of destination of the consignment of semen.

I.12:

Box

Seal number shall be indicated.

reference

I.19:

Box reference Total number of packages shall correspond to the number of containers.

I.26:

I.30:

Box reference "Type": Indicate semen.

"Species": Select amongst "Bos taurus", "Bison bison" or "Bubalus bubalis" as appropriate.

"Identification number": Indicate the identification number of each donor animal.

"Identification mark": Indicate the mark on the straw or other packages where the semen of the consignment is placed.

"Date of collection/production": Indicate the date on which the semen of consignment was collected.

"Approval or registration number of plant/establishment/centre": Indicate the unique approval number of the semen collection centre where the semen of the consignment was collected.

"Quantity": Indicate the number of straws or other packages with the same mark.

"Test": Indicate for BTV-test: point II.2.8.5. and/or point II.2.8.6, and/or for EHD-test: point II.2.9.4.1 and/or point II.2.9.4.2, if relevant.

Part II:

- Only semen collection centres approved by the competent authority and included in the register (1) referred to in Article 101(1), point (b), of Regulation (EU) 2016/429 of the European Parliament and of the Council and Article 7 of Delegated Regulation (EU) 2020/686.
- (2)Delete if not applicable.
- (3) Insert the name of the disease(s).
- (4) Insert the specific reference to the article(s), title, and number of the relevant legal act(s) adopted by the Commission providing for those requirements.
- (5)Insert the specific attestation(s) provided for in and required by the relevant legal act(s) adopted by the Commission, as referred to in Article 159(2), points (a), (b) and (c), of Regulation (EU) 2016/429.
- (6) Applicable for frozen semen.
- **(7)** Applicable for fresh and chilled semen.
- (8) Not applicable to animals which come from an establishment not free from enzootic bovine leukosis and which are less than 2 years of age as referred to in Article 20(2), point (a), of Delegated Regulation (EU) 2020/686.
- (9) Applicable only to seronegative animals.
- (10)Applicable only to bulls in semen production or having contact with bulls in semen production. Bulls returning to collection after a lay-off period of more than 6 months shall be tested during 30 days prior

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| | II. Health information | | | | | | |
| | to resuming production. | | 1 | | | | |
| | the commercial name of the | | | | | | |
| | semen diluent containing antibiotic(s). Certifying Officer/Official veterinarian | | | | | | |
| _ | Name (in capital letters) | Qualification and title | | | | | |
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