

Comparison of meteorological inputs for 2017 and 2018

Fig S1: Daily mean difference in wind speed multiplied by 2, along with the difference in aboveground DM and LAI for the model run using the parameterisation noted "calibration method 2", which used all available data.

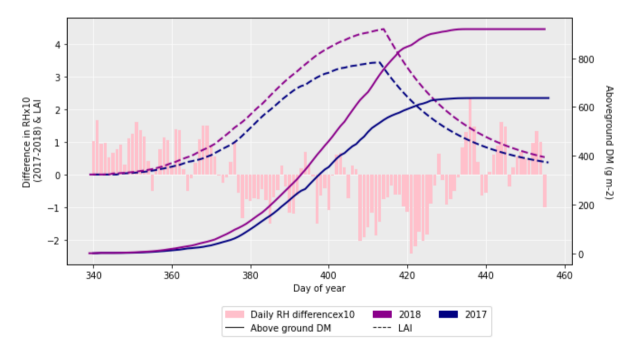


Fig. S2: Daily mean difference in RH multiplied by 10, along with the difference in aboveground DM and LAI for the model run using the parameterisation noted "calibration method 2", which used all available data.

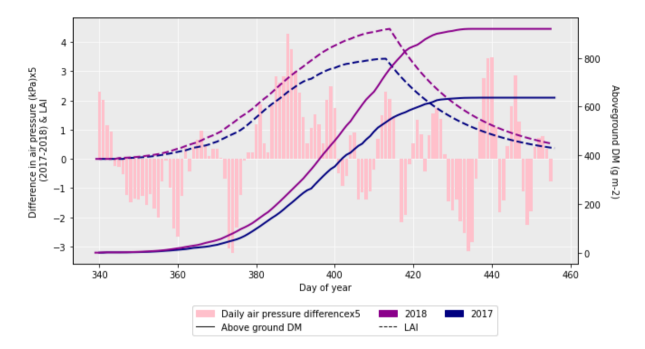


Fig. S3: Daily mean difference in air pressure multiplied by 5, along with the difference in aboveground DM and LAI for the model run using the parameterisation noted "calibration method 2", which used all available data.

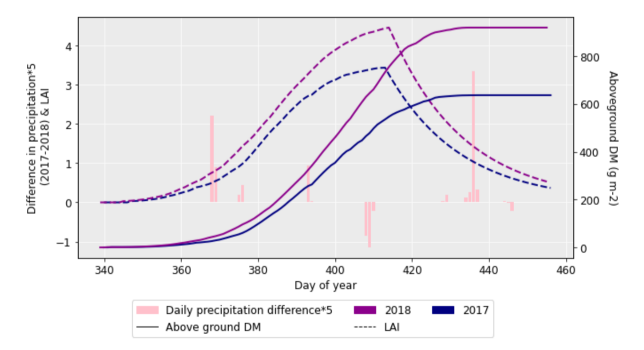
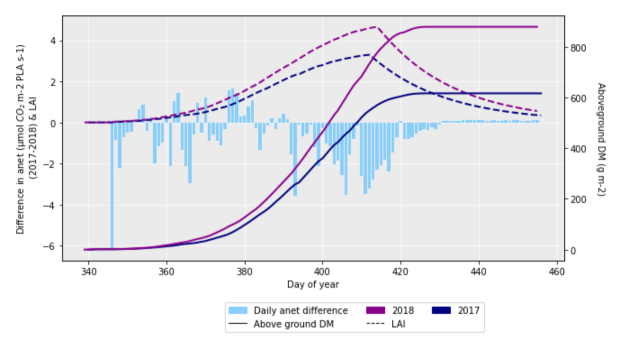


Fig. S4: Daily mean difference in precipitation multiplied by 5, along with the difference in aboveground DM and LAI for the model run using the parameterisation noted "calibration method 2", which used all available data. Wheat was assumed to be irrigated so lack of rain was not an issue in model runs



Comparison of photosynthetic processes

Fig. S5: The difference in net photosynthetic rate for 2017 and 2018 along with the difference in aboveground DM accumulation and LAI for the ambient treatment for the 2 years. The LAI and aboveground DM profiles are for the HD3118 cultivar.

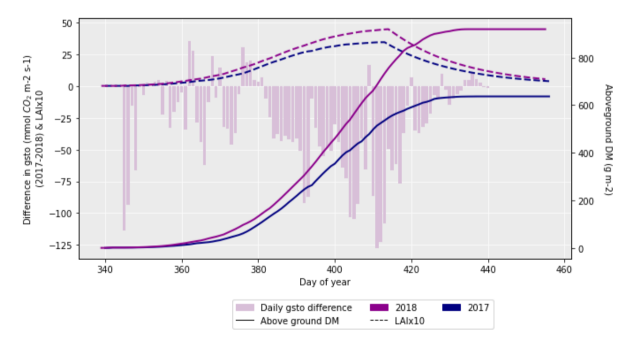


Fig. S6: The difference in sunlit stomatal conductance for 2017 and 2018 HUW234 cultivar along with the aboveground DM accumulation and LAI for both years. This run used the parameterisation of "calibration method 2" where all available data was used for calibration

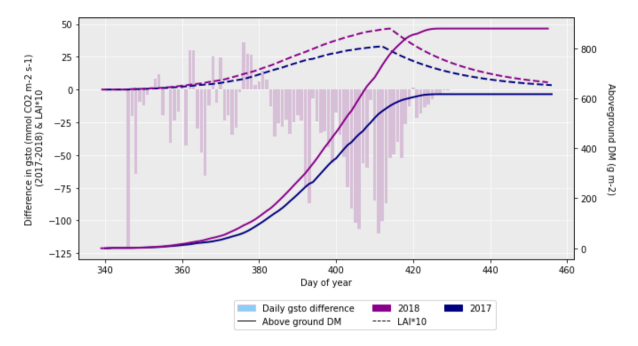


Fig. S7: The difference in sunlit stomatal conductance for 2017 and 2018 HD3118 cultivar along with the aboveground DM accumulation and LAI for both years. This run used the parameterisation of "calibration method 2" where all available data was used for calibration

Model calibration

Initially, the input data was split into 2 groups. The 2017 data was used to calibrate the model and the 2018 data was used to evaluate the model. However, with such limited data the 2017 calibration dataset was subject to overfitting and the parameterisation obtained in the calibration did not give good results for the 2018 evaluation dataset. The parameterisation using the 2017 data for calibration and 2018 for evaluation is referred to as calibration method 1, and the parameterisation used in the main body of the paper, which used all available data to develop a parameterisation, is referred to as calibration method 2.

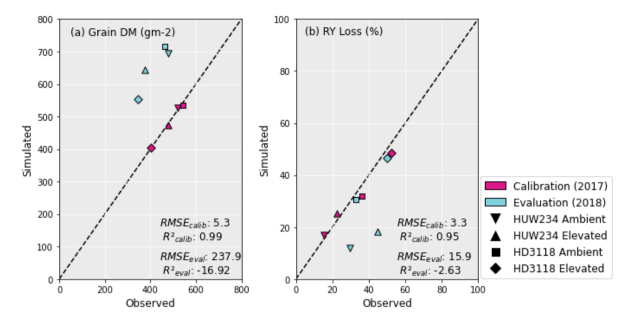


Figure S8: Calibration and evaluation of grain DM and RY loss using the DO₃SE-Crop model for the Varanasi dataset when using calibration method 1. RY loss was calculated comparative to preindustrial O₃ concentrations of 10 ppb (CLRTAP, 2017).

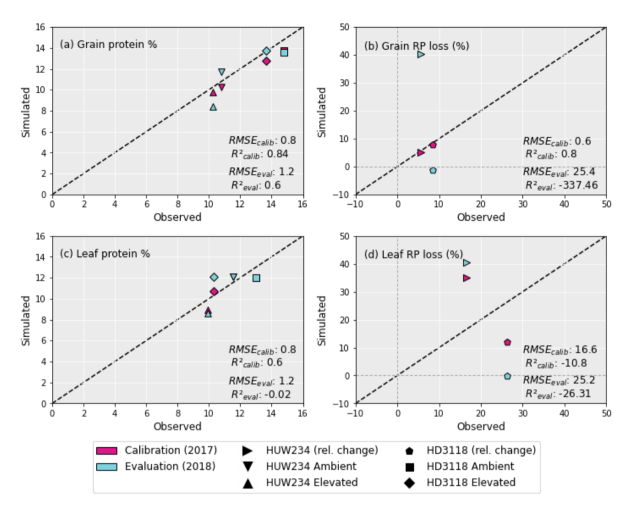


Fig. S9: Calibration and evaluation of the concentration of grain (a) and leaf (c) protein of HUW234 and HD3118 cultivars under ambient and elevated O_3 . Calibration and evaluation of the relative change in grain (b) and leaf (d) protein percentage. In figure (c) the ambient leaf protein % for the HUW234 and HD3118 cultivars in the calibration and evaluation were almost identical, hence the overlaid points. RMSE and R^2 of both the calibration and evaluation are indicated on the plot. These results use calibration method 1.

Correcting for the heating effect of the open top chamber

Data on the internal chamber and ambient air temperatures in Delhi, over the course of the wheat growing season in December 2018 to March 2019 were regressed against each other to obtain a regression with which to correct the input temperature data in the present study, as the air temperature sensor was external to the chambers.

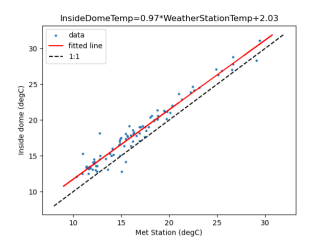


Figure S10: Regression between the air temperature as measured at the meteorological weather monitoring station and internal to the open top chambers in Delhi, during the wheat growth period 2018 to 2019. On average the open top chambers were approximately 2 degrees warmer than the ambient air.

Model parameterisation for calibration methods 1 and 2

Both the HUW234 and HD3118 cultivars were ran assuming the number of layers in the canopy was 4, and that there was 1 leaf population (nL=4, nP=1). The HUW234 cultivar parameterisation is given in Table S1, and the HD3118 cultivar parameterisation is given in Table S2.

Process	Parameter description	Calibrated Values		Unit
		Method 1	Method 2	
	Base temperature (T_b)	7	7	°C
	Optimum temperature (T _o)	25.99	25.99	°C
	Maximum temperature (T_m)	42.637	42.637	°C
Phenology	Plant emergence (<i>TT_{emr}</i>)	80	80	°C days
Filehology	Flag emergence (<i>TT_{flag,emr}</i>)	792	792	°C days
	Start anthesis (<i>TT_{astart}</i>)	1109	1109	°C days
	Mid-anthesis (<i>TT_{amid}</i>)	1181	1181	°C days
	Harvest (TT _{harv})	1668	1668	°C days
	Maximum carboxylation	99.3	99.3	$\mu mol \ CO_2 \ m^{-2} s^{-1}$
	capacity at 25 °C (V _{cmax,25})	00.0	00.0	$\mu m o c c o_2 m s$
	Leaf vertical N co-efficient (kN)	0	0	-
Photo-	Maximum rate of electron	138	138	$\mu mol \ CO_2 \ m^{-2} s^{-1}$
synthesis	transport at 25 °C ($J_{max,25}$)	100	100	$\mu m o c c o_2 m s$
	Parameter describing the			
	variation in relative stomatal	2.95	2.75	kPa
	conductance with VPD (VPD_0)			
	m (<i>m</i>)	7.4	7.4	-
Respiration	dark respiration (R_{dcoeff})	0.00972	0.00972	-
	growth respiration (R_g)	0.15	0.15	-

Table S1: The parameters that were calibrated for (changed from the default parameterisation) in DO₃SE-CropN Model for both calibration methods for the HUW234 cultivar

	Coefficient for determining DM partitioning (α_{root})	16.5	16.64	
	Coefficient for determining DM partitioning (β_{root})	-21	-20.5	-
	Coefficient for determining DM partitioning (α_{leaf})	20.477	18	-
DM	Coefficient for determining DM partitioning (eta_{leaf})	-24.5	-20	-
parameters	Coefficient for determining DM partitioning (α_{stem})	16.853	15.48	-
	Coefficient for determining DM partitioning (eta_{stem})	-17	-14.69	-
	Coefficient determining specific leaf area (Ω)	22.2	22.2	$m^2 kg^{-1}$
	Fraction of stem carbon in the reserve pool ($ au$)	0.75	0.75	-
	Fraction of DM in the harvest pool that goes to the grains (rest goes to the ear) (E_g)	0.85	0.85	-
	O₃ long term damage coefficient (γ3)	0.0000325	0.0000325	$(\mu mol \ O_3 \ m^{-2})^{-1}$
	O₃ long term damage coefficient determining senescence onset (γ4)	4.2553	3.1811	-
Ozone damage	O₃ long term damage coefficient determining maturity (γ5)	0.944	0.7742	-
	Critical accumulated stomatal O3 flux that determines the onset of leaf senescence (cL_{O_3})	8000	8500	mmol $O_3 m^{-2}$
	Pre-anthesis maximum N uptake (<i>NUP_{pre,max}</i>)	0.55	0.55	$g N m^{-2} da y^{-1}$
N uptake	Post-anthesis maximum N uptake (<i>NUP_{post,max}</i>)	0.3	0.3	$g N m^{-2} da y^{-1}$
Leaf and	Target leaf N concentration $([N_{leaf,target}])$	1	1	g N m ⁻² leaf area
stem N parameters	Target stem N concentration ([N _{stem,target}])	0.017	0.017	$N g^{-1} DW$
	\[¹ vstem,target])			

Grain N parameters	Ratio of N in grain to ear (f _{N,ear_grain})	0.95	0.95	-
	Alpha parameter controlling sigmoid N grain filling function (α_N)	23	23	-
	Beta parameter controlling sigmoid N grain filling function (β_N)	1.2	1.2	-
	Gradient of N remobilisation from the leaf under O_3 exposure (m_{leaf})	0.6	0.2	-
N re-	Intercept of N remobilisation from the leaf under O_3 exposure (c_{leaf})	10.89	10.89	-
mobilisation	Gradient of N remobilisation from the stem under O_3 exposure (m_{stem})	0.0325	0.0325	-
	Intercept of N remobilisation from the stem under O_3 exposure (c_{stem})	0.2293	0.2293	-
Antioxidant processes	Accumulated stomatal O ₃ flux above which N is only allocated to antioxidant pool (<i>fst_{end}</i>)	45000	45000	mmol $O_3 m^{-2}$
	Modifier to customise the O ₃ effect on antioxidants on the leaf (a_{leaf})	1	1	-
	Modifier to customise the O_3 effect on antioxidants on the stem (a_{stem})	2	2	-

Table S2: The parameters that were calibrated for (changed from the default parameterisation) in DO₃SE-CropN model for both calibration methods for the HD3118 cultivar

Process	Parameter description	Calibrate Method 1		Unit
Phenology	Base temperature (T_b)	6.992	6.992	°C
	Optimum temperature (T_o)	23	23	°C

	Maximum temperature (T_m)	43	43	°C
	Plant emergence (TT_{emr})	80	80	°C days
	Flag emergence (<i>TT_{flag,emr}</i>)	764	764	°C days
	Start anthesis (TT_{astart})	1050	1050	°C days
	Mid-anthesis (TT_{amid})	1093	1093	°C days
	Harvest (TT_{harv})	1450	1450	°C days
	Maximum carboxylation			
	capacity at 25 °C ($V_{cmax,25}$)	101.6	101.6	$\mu mol \ CO_2 \ m^{-2} s^{-1}$
	Leaf vertical N co-efficient (kN)	0	0.2	-
	Maximum rate of electron			1 - 2 - 1
Photo-	transport at 25 °C ($J_{max,25}$)	144	144	$\mu mol \ CO_2 \ m^{-2} s^{-1}$
synthesis	Parameter describing the			
	variation in relative stomatal	3.85	3	kPa
	conductance with VPD (VPD_0)			
	m (<i>m</i>)	8.1	7.586	-
Despiration	dark respiration (R_{dcoeff})	0.00726	0.00726	-
Respiration	growth respiration (R_g)	0.2	0.15	-
	Coefficient for determining	16	16	
	DM partitioning (α_{root})	10	10	-
	Coefficient for determining	-21.5	-20.5	
	DM partitioning (eta_{root})	-21.5	-20.5	-
	Coefficient for determining	17.5	18	_
	DM partitioning (α_{leaf})	17.5	10	-
	Coefficient for determining	-19.921	-20	
DM	DM partitioning (eta_{leaf})	-10.021	-20	_
parameters	Coefficient for determining	15.15	16.6	_
	DM partitioning (α_{stem})	10.10	10.0	
	Coefficient for determining	-15.714	-16.5	_
	DM partitioning (eta_{stem})			
	Coefficient determining	22.2	22.2	$m^2 kg^{-1}$
	specific leaf area (Ω)			
	Fraction of stem carbon in	0.7	0.7	-
	the reserve pool (τ)			
	Fraction of DM in the			
	harvest pool that goes to	0.85	0.85	-
	the grains (rest goes to the ear) (E_a)			
	O_3 long term damage			
	coefficient (γ 3)	0.00008	0.0000377	' $(\mu mol \ O_3 \ m^{-2})^{-1}$
	O_3 long term damage			
Ozone damage	coefficient determining	0.9938	6.81	_
	senescence onset (γ 4)	0.0000	0.01	
	O_3 long term damage			
	coefficient determining	0.92	1.4	-
	maturity (γ 5)			

	Critical accumulated stomatal O3 flux that determines the onset of leaf senescence (cL_{O_3})	12000	8500	mmol $O_3 m^{-2}$
Nuntoko	Pre-anthesis maximum N uptake (<i>NUP_{pre,max}</i>)	0.65	0.5	$g N m^{-2} da y^{-1}$
N uptake	Post-anthesis maximum N uptake (<i>NUP_{post,max}</i>)	0.4	0.2	$g N m^{-2} da y^{-1}$
Leaf and stem N	Target leaf N concentration ([N _{leaf,target}])	1	1.2	g N m ⁻² leaf area
parameters	Target stem N concentration ([N _{stem,target}])	0.025	0.02	$N g^{-1} DW$
	Ratio of N in grain to ear (f _{N,ear_grain})	0.95	0.95	-
Grain N parameters	Alpha parameter controlling sigmoid N grain filling function (α_N)	23	23	-
	Beta parameter controlling sigmoid N grain filling function (β_N)	1.2	1.2	-
	Gradient of N remobilisation from the leaf under O_3 exposure (m_{leaf})	0.2	0.2	-
N re- mobilisation	Intercept of N remobilisation from the leaf under O_3 exposure (c_{leaf})	10.89	10.89	-
	Gradient of N remobilisation from the stem under O_3 exposure (m_{stem})	0.2293	0.0335	-
	Intercept of N remobilisation from the stem under O_3 exposure (c_{stem})	0.03425	0.15	-

	Accumulated stomatal O_3 flux above which N is only allocated to antioxidant pool (fst_{end})	35000	75000	mmol $O_3 m^{-2}$
Antioxidant processes	Modifier to customise the O ₃ effect on antioxidants on the leaf (a _{leaf})	0.6	1	-
	Modifier to customise the O_3 effect on antioxidants on the stem (a_{stem})	10	2	-

Senescence

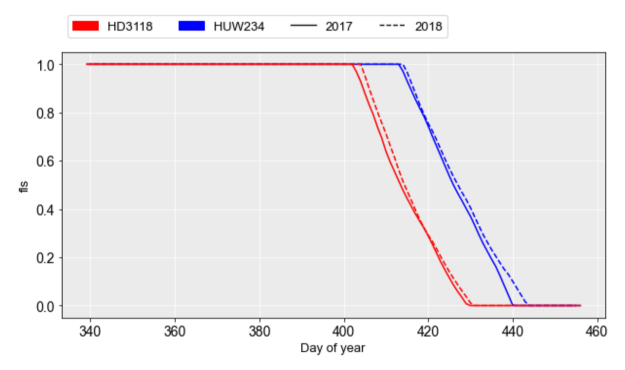


Figure S11: Graph of fls, the factor describing leaf senescence, where 0 is full senescence and 1 is no senescence, for both cultivars and years